

# Package ‘methyICC’

July 8, 2025

**Title** Estimate the cell composition of whole blood in DNA methylation samples

**Version** 1.23.0

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**Depends** R (>= 3.6), FlowSorted.Blood.450k

**Suggests** rmarkdown, knitr, testthat (>= 2.1.0), BiocGenerics, BiocStyle, tidyrr, ggplot2

**Description** A tool to estimate the cell composition of DNA methylation whole blood sample measured on any platform technology (microarray and sequencing).

**biocViews** Microarray, Sequencing, DNAMethylation, MethylationArray, MethylSeq, WholeGenome

**VignetteBuilder** knitr

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|                                |                         |
|--------------------------------|-------------------------|
| <code>.extract_raw_data</code> | <i>Extract raw data</i> |
|--------------------------------|-------------------------|

---

## Description

Extract the methylation values and GRanges objects

## Usage

```
.extract_raw_data(object)
```

## Arguments

`object` an object can be a `RGChannelSet`, `GenomicMethylSet` or `BSseq` object

## Value

A list preprocessed objects from the `RGChannelSet`, `GenomicMethylSet` or `BSseq` objects to be used in `.preprocess_estimatecc()`.

---

.find\_dmrs*Finding differentially methylated regions*

---

**Description**

This function uses the FlowSorted.Blood.450k whole blood reference methylomes with six cell types to identify differentially methylated regions.

**Usage**

```
.find_dmrs(verbose = TRUE, gr_target = NULL, include_cpgs = FALSE,
  include_dmrs = TRUE, num_cpgs = 50, num_regions = 50,
  bumphunter_beta_cutoff = 0.2, dmr_up_cutoff = 0.5,
  dmr_down_cutoff = 0.4, dmr_pval_cutoff = 1e-11,
  cpg_pval_cutoff = 1e-08, cpg_up_dm_cutoff = 0,
  cpg_down_dm_cutoff = 0, pairwise_comparison = FALSE,
  mset_train_flow_sort = NULL)
```

**Arguments**

|                        |                                                                                                                                                                                                                                                                                     |
|------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| verbose                | TRUE/FALSE argument specifying if verbose messages should be returned or not. Default is TRUE.                                                                                                                                                                                      |
| gr_target              | Default is NULL. However, the user can provide a GRanges object from the object in estimatecc. Before starting the procedure to find differentially methylated regions, the intersection of the gr_target and GRanges object from the reference methylomes (FlowSorted.Blood.450k). |
| include_cpgs           | TRUE/FALSE. Should individual CpGs be returned. Default is FALSE.                                                                                                                                                                                                                   |
| include_dmrs           | TRUE/FALSE. Should differentially methylated regions be returned. Default is TRUE. User can turn this to FALSE and search for only CpGs.                                                                                                                                            |
| num_cpgs               | The max number of CpGs to return for each cell type. Default is 50.                                                                                                                                                                                                                 |
| num_regions            | The max number of DMRs to return for each cell type. Default is 50.                                                                                                                                                                                                                 |
| bumphunter_beta_cutoff | The cutoff threshold in bumphunter() in the bumphunter package.                                                                                                                                                                                                                     |
| dmr_up_cutoff          | A cutoff threshold for identifying DMRs that are methylated in one cell type, but not in the other cell types.                                                                                                                                                                      |
| dmr_down_cutoff        | A cutoff threshold for identifying DMRs that are not methylated in one cell type, but methylated in the other cell types.                                                                                                                                                           |
| dmr_pval_cutoff        | A cutoff threshold for the p-values when identifying DMRs that are methylated in one cell type, but not in the other cell types (or vice versa).                                                                                                                                    |
| cpg_pval_cutoff        | A cutoff threshold for the p-values when identifying differentially methylated CpGs that are methylated in one cell type, but not in the other cell types (or vice versa).                                                                                                          |
| cpg_up_dm_cutoff       | A cutoff threshold for identifying differentially methylated CpGs that are methylated in one cell type, but not in the other cell types.                                                                                                                                            |

|                                   |                                                                                                                                                                                                                |
|-----------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>cpg_down_dm_cutoff</code>   | A cutoff threshold for identifying differentially methylated CpGs that are not methylated in one cell type, but are methylated in the other cell types.                                                        |
| <code>pairwise_comparison</code>  | TRUE/FAISE of whether all pairwise comparisons (e.g. methylated in Granulocytes and Monocytes, but not methylated in other cell types). Default if FALSE.                                                      |
| <code>mset_train_flow_sort</code> | Default is NULL. However, a user can provide a <code>MethylSet</code> object after processing the <code>FlowSorted.Blood.450k</code> dataset. The default normalization is <code>preprocessIllumina()</code> . |

**Value**

A list of data frames and `GRanges` objects.

---

|                              |                        |
|------------------------------|------------------------|
| <code>.initializeMLEs</code> | <i>.initializeMLEs</i> |
|------------------------------|------------------------|

---

**Description**

Helper functions to initialize MLEs in `estimatecc()`.

**Usage**

```
.initializeMLEs(init_param_method, n, K, Ys, Zs, a0init, a1init, sig0init,
  sig1init, tauinit)
```

**Arguments**

|                                |                                                                                                                                                                                                                                               |
|--------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>init_param_method</code> | method to initialize parameter estimates. Choose between "random" (randomly sample) or "known_regions" (uses unmethylated and methylated regions that were identified based on Reinus et al. (2012) cell sorted data.). Defaults to "random". |
| <code>n</code>                 | Number of samples                                                                                                                                                                                                                             |
| <code>K</code>                 | Number of cell types                                                                                                                                                                                                                          |
| <code>Ys</code>                | observed methylation levels in samples provided by user of dimension $R \times n$                                                                                                                                                             |
| <code>Zs</code>                | Cell type specific regions of dimension $R \times K$                                                                                                                                                                                          |
| <code>a0init</code>            | Default NULL. Initial mean methylation level in unmethylated regions                                                                                                                                                                          |
| <code>a1init</code>            | Default NULL. Initial mean methylation level in methylated regions                                                                                                                                                                            |
| <code>sig0init</code>          | Default NULL. Initial var methylation level in unmethylated regions                                                                                                                                                                           |
| <code>sig1init</code>          | Default NULL. Initial var methylation level in methylated regions                                                                                                                                                                             |
| <code>tauint</code>            | Default NULL. Initial var for measurement error                                                                                                                                                                                               |

**Value**

A list of MLE estimates to be used in `estimatecc()`.

---

|                          |                          |
|--------------------------|--------------------------|
| <i>.initialize_theta</i> | <i>.initialize_theta</i> |
|--------------------------|--------------------------|

---

**Description**

Creates a container with initial theta parameter estimates

**Usage**

```
.initialize_theta(n, K, alpha0 = NULL, alpha1 = NULL, sig0 = NULL,
  sig1 = NULL, tau = NULL)
```

**Arguments**

|        |                                                                      |
|--------|----------------------------------------------------------------------|
| n      | Number of samples                                                    |
| K      | Number of cell types                                                 |
| alpha0 | Default NULL. Initial mean methylation level in unmethylated regions |
| alpha1 | Default NULL. Initial mean methylation level in methylated regions   |
| sig0   | Default NULL. Initial var methylation level in unmethylated regions  |
| sig1   | Default NULL. Initial var methylation level in methylated regions    |
| tau    | Default NULL. Initial var for measurement error                      |

**Value**

A data frame with initial parameter estimates to be used in *.initializeMLEs()*.

---

|                         |                         |
|-------------------------|-------------------------|
| <i>.methylcc_engine</i> | <i>.methylcc_engine</i> |
|-------------------------|-------------------------|

---

**Description**

Helper function for *estimatecc*

**Usage**

```
.methylcc_engine(Ys, Zs, current_pi_mle, current_theta, epsilon, max_iter)
```

**Arguments**

|                |                                                                            |
|----------------|----------------------------------------------------------------------------|
| Ys             | observed methylation levels in samples provided by user of dimension R x n |
| Zs             | Cell type specific regions of dimension R x K                              |
| current_pi_mle | cell composition MLE estimates of dimension K x n                          |
| current_theta  | other parameter estimates in EM algorithm                                  |
| epsilon        | Add here.                                                                  |
| max_iter       | Add here.                                                                  |

**Value**

A list of MLE estimates that is used in *estimatecc()*.

---

|                  |                         |
|------------------|-------------------------|
| .methyllcc_estep | <i>Expectation step</i> |
|------------------|-------------------------|

---

### Description

Expectation step in EM algorithm for methyllCC

### Usage

```
.methyllcc_estep(Ys, Zs, current_pi_mle, current_theta, meth_status = 0)
```

### Arguments

|                |                                                                                                                 |
|----------------|-----------------------------------------------------------------------------------------------------------------|
| Ys             | observed methylation levels in samples provided by user of dimension R x n                                      |
| Zs             | Cell type specific regions of dimension R x K                                                                   |
| current_pi_mle | cell composition MLE estimates of dimension K x n                                                               |
| current_theta  | other parameter estimates in EM algorithm                                                                       |
| meth_status    | Indicator function corresponding to regions that are unmethylated (meth_status=0) or methylated (meth_status=1) |

### Value

List of expected value of the first two moments of the random effects (or the E-Step in the EM algorithm) used in .methyllcc\_engine()

---

|                  |                          |
|------------------|--------------------------|
| .methyllcc_mstep | <i>Maximization step</i> |
|------------------|--------------------------|

---

### Description

Maximization step in EM Algorithm for methyllCC

### Usage

```
.methyllcc_mstep(Ys, Zs, current_pi_mle, current_theta, estep0, estep1)
```

### Arguments

|                |                                                                            |
|----------------|----------------------------------------------------------------------------|
| Ys             | observed methylation levels in samples provided by user of dimension R x n |
| Zs             | Cell type specific regions of dimension R x K                              |
| current_pi_mle | cell composition MLE estimates of dimension K x n                          |
| current_theta  | other parameter estimates in EM algorithm                                  |
| estep0         | Results from expectation step for unmethylated regions                     |
| estep1         | Results from expectation step for methylated regions                       |

### Value

A list of the updated MLEs (or the M-Step in the EM algorithm) used in .methyllcc\_engine()

---

*.pick\_target\_positions**Pick target positions*

---

**Description**

Pick probes from target data using the indices in `dmp_regions`

**Usage**

```
.pick_target_positions(target_granges, target_object = NULL,
  target_cvg = NULL, dmp_regions)
```

**Arguments**

|                             |                                                                                                                                                                                                       |
|-----------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>target_granges</code> | add more here.                                                                                                                                                                                        |
| <code>target_object</code>  | an optional argument which contains the meta-data for <code>target_granges</code> . If <code>target_granges</code> already contains the meta-data, do not need to supply <code>target_object</code> . |
| <code>target_cvg</code>     | coverage reads for the target object                                                                                                                                                                  |
| <code>dmp_regions</code>    | differentially methylated regions                                                                                                                                                                     |

**Value**

A list of `GRanges` objects to be used in `.preprocess_estimatecc()`

---

*.preprocess\_estimatecc**.preprocess\_estimatecc*

---

**Description**

This function preprocesses the data before the `estimatecc()` function

**Usage**

```
.preprocess_estimatecc(object, verbose = TRUE,
  init_param_method = "random",
  celltype_specific_dmrs = celltype_specific_dmrs)
```

**Arguments**

|                                     |                                                                                                                                                                                                                                               |
|-------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>object</code>                 | an object can be a <code>RGChannelSet</code> , <code>GenomicMethylSet</code> or <code>BSseq</code> object                                                                                                                                     |
| <code>verbose</code>                | TRUE/FALSE argument specifying if verbose messages should be returned or not. Default is TRUE.                                                                                                                                                |
| <code>init_param_method</code>      | method to initialize parameter estimates. Choose between "random" (randomly sample) or "known_regions" (uses unmethylated and methylated regions that were identified based on Reinus et al. (2012) cell sorted data.). Defaults to "random". |
| <code>celltype_specific_dmrs</code> | cell type specific differentially methylated regions (DMRs).                                                                                                                                                                                  |

**Value**

A list of object to be used in estimatecc

---

|                       |                 |
|-----------------------|-----------------|
| <code>.splitit</code> | <i>.splitit</i> |
|-----------------------|-----------------|

---

**Description**

helper function to split along a variable

**Usage**

```
.splitit(x)
```

**Arguments**

`x`                      a vector

**Value**

A list to be used in find\_dmrs()

---

|                    |                                                                                |
|--------------------|--------------------------------------------------------------------------------|
| <code>.WFun</code> | <i>Helper function to take the product of Z and cell composition estimates</i> |
|--------------------|--------------------------------------------------------------------------------|

---

**Description**

Helper function which is the product of Z and pi\_mle

**Usage**

```
.WFun(Zs, pi_mle)
```

**Arguments**

`Zs`                      Cell type specific regions of dimension R x K  
`pi_mle`                   cell composition MLE estimates

**Value**

A list of output after taking the product of Z and cell composition mle estimates to be used in `.methyloc_estep()`.



---

|             |                                                                     |
|-------------|---------------------------------------------------------------------|
| cell_counts | <i>Generic function that returns the cell composition estimates</i> |
|-------------|---------------------------------------------------------------------|

---

### Description

Given a estimatecc object, this function returns the cell composition estimates

Accessors for the 'cell\_counts' slot of a estimatecc object.

### Usage

```
cell_counts(object)

## S4 method for signature 'estimatecc'
cell_counts(object)
```

### Arguments

object            an object of class estimatecc.

### Value

Returns the cell composition estimates

### Examples

```
# This is a reduced version of the FlowSorted.Blood.450k
# dataset available by using BiocManager::install("FlowSorted.Blood.450k"),
# but for purposes of the example, we use the smaller version
# and we set \code{demo=TRUE}. For any case outside of this example for
# the package, you should set \code{demo=FALSE} (the default).

dir <- system.file("data", package="methy1CC")
files <- file.path(dir, "FlowSorted.Blood.450k.sub.RData")
if(file.exists(files)){
  load(file = files)

  set.seed(12345)
  est <- estimatecc(object = FlowSorted.Blood.450k.sub, demo = TRUE)
  cell_counts(est)
}
```

---

|            |                                                 |
|------------|-------------------------------------------------|
| estimatecc | <i>Estimate cell composition from DNAm data</i> |
|------------|-------------------------------------------------|

---

### Description

Estimate cell composition from DNAm data

## Usage

```
estimatecc(object, find_dmrs_object = NULL, verbose = TRUE,
  epsilon = 0.01, max_iter = 100, take_intersection = FALSE,
  include_cpgs = FALSE, include_dmrs = TRUE,
  init_param_method = "random", a0init = NULL, a1init = NULL,
  sig0init = NULL, sig1init = NULL, tauinit = NULL, demo = FALSE)
```

## Arguments

|                                |                                                                                                                                                                                                                                               |
|--------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>object</code>            | an object can be a <code>RGChannelSet</code> , <code>GenomicMethylSet</code> or <code>BSseq</code> object                                                                                                                                     |
| <code>find_dmrs_object</code>  | If the user would like to supply different differentially methylated regions, they can use the output from the <code>find_dmrs</code> function to supply different regions to <code>estimatecc</code> .                                       |
| <code>verbose</code>           | TRUE/FALSE argument specifying if verbose messages should be returned or not. Default is TRUE.                                                                                                                                                |
| <code>epsilon</code>           | Threshold for EM algorithm to check for convergence. Default is 0.01.                                                                                                                                                                         |
| <code>max_iter</code>          | Maximum number of iterations for EM algorithm. Default is 100 iterations.                                                                                                                                                                     |
| <code>take_intersection</code> | TRUE/FALSE asking if only the CpGs included in object should be used to find DMRs. Default is FALSE.                                                                                                                                          |
| <code>include_cpgs</code>      | TRUE/FALSE. Should individual CpGs be returned. Default is FALSE.                                                                                                                                                                             |
| <code>include_dmrs</code>      | TRUE/FALSE. Should differentially methylated regions be returned. Default is TRUE.                                                                                                                                                            |
| <code>init_param_method</code> | method to initialize parameter estimates. Choose between "random" (randomly sample) or "known_regions" (uses unmethylated and methylated regions that were identified based on Reinus et al. (2012) cell sorted data.). Defaults to "random". |
| <code>a0init</code>            | Default NULL. Initial mean methylation level in unmethylated regions                                                                                                                                                                          |
| <code>a1init</code>            | Default NULL. Initial mean methylation level in methylated regions                                                                                                                                                                            |
| <code>sig0init</code>          | Default NULL. Initial var methylation level in unmethylated regions                                                                                                                                                                           |
| <code>sig1init</code>          | Default NULL. Initial var methylation level in methylated regions                                                                                                                                                                             |
| <code>tauint</code>            | Default NULL. Initial var for measurement error                                                                                                                                                                                               |
| <code>demo</code>              | TRUE/FALSE. Should the function be used in demo mode to shorten examples in package. Defaults to FALSE.                                                                                                                                       |

## Value

A object of the class `estimatecc` that contains information about the cell composition estimation (in the `summary` slot) and the cell composition estimates themselves (in the `cell_counts` slot).

## Examples

```
# This is a reduced version of the FlowSorted.Blood.450k
# dataset available by using BiocManager::install("FlowSorted.Blood.450k"),
# but for purposes of the example, we use the smaller version
# and we set \code{demo=TRUE}. For any case outside of this example for
# the package, you should set \code{demo=FALSE} (the default).
```

```

dir <- system.file("data", package="methyLCC")
files <- file.path(dir, "FlowSorted.Blood.450k.sub.RData")
if(file.exists(files)){
  load(file = files)

  set.seed(12345)
  est <- estimatecc(object = FlowSorted.Blood.450k.sub, demo = TRUE)
  cell_counts(est)
}

```

---

|                  |                             |
|------------------|-----------------------------|
| estimatecc-class | <i>the estimatecc class</i> |
|------------------|-----------------------------|

---

## Description

Objects of this class store all the values needed information to work with a estimatecc object

## Value

summary returns the summary information about the cell composition estimate procedure and cell\_counts returns the cell composition estimates

## Slots

summary information about the samples and regions used to estimate cell composition  
 cell\_counts cell composition estimates

## Examples

```

# This is a reduced version of the FlowSorted.Blood.450k
# dataset available by using BiocManager::install("FlowSorted.Blood.450k"),
# but for purposes of the example, we use the smaller version
# and we set \code{demo=TRUE}. For any case outside of this example for
# the package, you should set \code{demo=FALSE} (the default).

dir <- system.file("data", package="methyLCC")
files <- file.path(dir, "FlowSorted.Blood.450k.sub.RData")
if(file.exists(files)){
  load(file = files)

  set.seed(12345)
  est <- estimatecc(object = FlowSorted.Blood.450k.sub, demo = TRUE)
  cell_counts(est)
}

```

---

FlowSorted.Blood.450k.sub

*A reduced size of the FlowSorted.Blood.450k dataset*

---

### Description

A reduced size of the FlowSorted.Blood.450k dataset

The object was created using the script in /inst and located in the /data folder.

### Format

A RGset object with 2e5 rows (probes) and 6 columns (whole blood samples).

---

offMethRegions

*Unmethylated regions for all celltypes*

---

### Description

This is the script used to create the offMethRegions data set. The purpose is use in the estimate\_cc() function.

The object was created using the script in /inst and located in the /data folder.

### Format

add more here.

---

onMethRegions

*Methylated regions for all celltypes*

---

### Description

This is the script used to create the onMethRegions data set. The purpose is use in the estimate\_cc() function.

The object was created using the script in /inst and located in the /data folder.

### Format

add more here.

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