

# MBCB

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MBCBExpressionData *MBCB - Bayesian Background Correction for Illumina Beadarray*

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## Description

The MBCBExpressionData dataset consists of two dataframes, expressionSignal and negativeControl. See the vignette for more detail regarding these data.

## Usage

```
data(MBCBExpressionData)
```

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bg.mcmc *MBCB - Bayesian Background Correction for Illumina Beadarray*

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## Description

This function provides the means of using only the MCMC (Bayesian) background correction method for the Illumina platform.

## Usage

```
bg.mcmc(iter=500, burn=200)
```

## Arguments

`iter` The iteration count for the Bayesian correction.  
`burn` The number of iterations to burn for the Bayesian correction.

## Value

This function returns an array of alpha, mu, and sigma values representing the values computed during the mcmc trial.

**Note**

This function makes use of two global variables. It will expect `obsbead` and `obsnc` are both established prior to calling this function. Obviously, this is not ideal, but R's pass-by-value functionality hindered the ability to pass these matrices as parameters. Using global variables increases performance substantially.

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**See Also**

[mbcb.main](#)

**Examples**

```
data(MBCBExpressionData)
# Use of global variables is obviously not ideal, but with R's pass-by-value
# setup, we quickly run out of memory without using them on such large
# arrays

#all of the signals from sample #2
obsbead <- expressionSignal[,2]

#the negative control values for this sample
obsnc <- negativeControl[,2]

#compute the alpha, mu, and sigma values
bg.mcmc();
```

---

bg.rma

*MBCB - Robust Multi-Array Average Background Correction for Illumina Beadarray*

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**Description**

This function provides the means of using only the Robust Multi-Array Average background correction method for the Illumina platform.

**Usage**

```
bg.rma(pm, n.pts = 2^14)
```

**Arguments**

<code>pm</code>	The data to be background corrected.
<code>n.pts</code>	Pertains to the specificity or accuracy of the method. $2^{14}$ is the default.

**Value**

A list with two values:

**ex.rma:** A vector containing the background-corrected values.

**para:** The statistical summary of the computation.

**Note**

This method does not make use of negative control beads.

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**See Also**

[mbcb.main](#)

**Examples**

```
data(MBCBExpressionData)

bg.rma(expressionSignal[,2]);
```

---

mbcb.correct

*MBCB - Model-Based Background Correction for Illumina Beadarray*

---

**Description**

This function is used to background-correct the provided data using the selected correction methods. Normalization is *not* applied.

**Usage**

```
mbcb.correct (g,
  control,
  npBool=TRUE,
  rmaBool=FALSE,
  mleBool=FALSE,
  bayesBool=FALSE,
  gmleBool=FALSE,
  iter=500,
  burn=200,
  isRawBead=FALSE)
```

**Arguments**

<code>g</code>	The data representing the signal file.
<code>control</code>	The data representing the control file.
<code>npBool</code>	A boolean value representing the desire to compute the non-parametric background correction values. Set to true if you wish to compute Non-Parametric background correction.
<code>rmaBool</code>	A boolean value representing the desire to compute the RMA background correction values. Set to true if you wish to compute RMA background correction.
<code>mleBool</code>	A boolean value representing the desire to compute the MLE background correction values. Set to true if you wish to compute MLE background correction.
<code>bayesBool</code>	A boolean value representing the desire to compute the Bayes background correction values. Set to true if you wish to compute Bayes background correction.
<code>gmleBool</code>	A boolean value representing the desire to compute the GMLE background correction values. Set to true if you wish to compute GMLE background correction.
<code>iter</code>	The iteration count; only used in Bayesian correction.
<code>burn</code>	The number of iterations which will be burned; only used in Bayesian correction.
<code>isRawBead</code>	A boolean value representing whether the input files are bead-level or bead-type. If the input is bead-level, set this value to True so that the raw bead-level values can be summarized to bead-type data.

**Value**

This function returns a complex list which can be grouped into two categories:  
Background-corrected Values:

**NP:** The background corrected values of the Non-Parametric method (or an empty data.frame if this method was not used).

**RMA:** The background corrected values of the RMA method (or an empty data.frame if this method was not used).

**MLE:** The background corrected values of the MLE method (or an empty data.frame if this method was not used).

**Bayes:** The background corrected values of the Bayesian method (or an empty data.frame if this method was not used).

## Average Values

**AvgNP:** A data.frame of statistics pertaining to the average of the NP normalization method (or an empty data.frame if this method was not used).

**AvgRMA:** A data.frame of statistics pertaining to the average of the RMA method (or an empty data.frame if this method was not used).

**AvgMLE:** A data.frame of statistics pertaining to the average of the MLE method (or an empty data.frame if this method was not used).

**AvgBayes:** A data.frame of statistics pertaining to the average of the Bayesian method (or an empty data.frame if this method was not used).

These values have not been normalized or log<sub>2</sub> transformed. See [mbcb.main](#) for such functionality.

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**See Also**

[mbcb.main](#)

**Examples**

```
data(MBCBExpressionData)
mbcb.correct(expressionSignal, negativeControl);
```

---

mbcb.gui

*MBCB - Model-Based Background Correction for Illumina Beadarray*

---

**Description**

This file provides the Graphical-User-Interface for the MBCB package.

**Usage**

```
mbcb.gui()
```

**Value**

This GUI will allow you to easily input data files and a negative-control file and will output a log2-transformed background-corrected file.

**Note**

The input files should be tab-delimited files in the following form:

*signal.txt*

```
g.1 g.2 g.3 g.4
10181072_239_rc-S 160.3 776.4 135.8 407.7
10181072_290-S 138.1 219.8 122.1 142.4
... ..
```

*neg-con.txt*

```
id g.1 g.2 g.3 g.4
50133 127 213.5 82 103
50315 232 295 143.5 156
... ..
```

Note that both of these examples come from a test involving four trials. Your data could use whatever number you'd like.

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**See Also**

[mbcb.main](#)

**Examples**

```
data(MBCBExpressionData)

#create files from the data provided in this package in the current directory
write.table(expressionSignal, 'signal.txt', sep="\t");
write.table(negativeControl, 'negative.control.txt', sep="\t");

#open the GUI; you can use the files just created as input for the signal and
# negative control files.
mbcb.gui();
```

---

mbcb.main

*MBCB - Model-Based Background Correction for Illumina Beadarray*

---

**Description**

This is the main function which incorporates all the others. This should be the most straightforward and autonomous function in the MBCB package.

**Usage**

```
mbcb.main (signal,
           control,
           npBool=TRUE,
           rmaBool=FALSE,
           mleBool=FALSE,
           bayesBool=FALSE,
           gmleBool=FALSE,
           paramEstFile="param-est",
           bgCorrectedFile="bgCorrected",
           iter=500,
           burn=200,
           normMethod="none",
           isRawBead=FALSE)
```

**Arguments**

signal	The data representing the signal file.
control	The data representing the control file.
npBool	A boolean value representing the desire to compute the non-parametric background correction values.

rmaBool	A boolean value representing the desire to compute the RMA background correction values.
mleBool	A boolean value representing the desire to compute the MLE background correction values.
bayesBool	A boolean value representing the desire to compute the Bayes background correction values.
gmleBool	A boolean value representing the desire to compute the GMLE background correction values.
paramEstFile	The <i>base</i> file name to which suffixes and a file extension will be appended (i.e. 'C:/output'). These files will store the parameter estimates of each background correction method selected.
bgCorrectedFile	The <i>base</i> file name to which suffixes and a file extension will be appended (i.e. 'C:/output'). These files will store the background corrected intensities
iter	The iteration count; only used in Bayesian correction.
burn	The number of iterations which will be burned; only used in Bayesian correction.
normMethod	The normalization method to be used. By default, none will be applied. The choices are: <i>none</i> - no normalization will be applied. <i>quant</i> - Quantile-Quantile normalization will be applied (requires the <i>affy</i> and <i>affyio</i> packages be present). <i>median</i> - Median or Global normalization will be applied.
isRawBead	A boolean value representing whether the input files are bead-level or bead-type. If the input is bead-level, set this value to True so that the raw bead-level values can be summarized to bead-type data.

### Value

The function will compute and output log<sub>2</sub>-transformed values for the desired background correction methods. Rather than returning this data as R objects, the output is written to files based on the names given in *paramEstFile* and *bgCorrectedFile*.

### Note

You can use [mbcb.parseFile](#) to create the signal and control matrices from the given files.

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### See Also

[mbcb.correct](#) [mbcb.parseFile](#)

### Examples

```
data(MBCBExpressionData)
#Use of global variables is obviously not ideal, but with R's pass-by-value
# setup, we quickly run out of memory without using them on such large
```

```
# arrays
mbcb.main(expressionSignal, negativeControl);
```

---

```
mbcb.parseFile      MBCB - Model-Based Background Correction for Illumina Beadarray
```

---

## Description

This function is used to read the files provided into a format which will be usable by the MBCB package (data.frame).

## Usage

```
mbcb.parseFile(sigFile, conFile, isRawBead = FALSE)
```

## Arguments

sigFile	The file-name (character string) representing the signal file.
conFile	The file-name (character string) representing the control file.
isRawBead	A boolean value representing whether the input files are bead-level or bead-type. If the input is bead-level, set this value to True so that the raw bead-level values can be summarized to bead-type data.

## Value

This function will return a list containing two data frames: *sig* and *con*.

**sig:** Represents the signal file.

**con:** Represents the negative control data.frame.

## Note

The input files should be tab-delimited files in the following form:

*signal.txt*

```
g.1 g.2 g.3 g.4
10181072_239_rc-S 160.3 776.4 135.8 407.7
10181072_290-S 138.1 219.8 122.1 142.4
... ..
```

*neg-con.txt*

```
id g.1 g.2 g.3 g.4
50133 127 213.5 82 103
50315 232 295 143.5 156
... ..
```

Note that both of these examples come from a test involving four trials. Your data could use whatever number you'd like.



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**See Also**

[mbcb.correct](#)

**Examples**

```
data(MBCBExpressionData)

# Create files from the data provided in this package in the current
#   directory
# Obviously, this is the opposite of what the function does, but we need to
#   write sample files for the sake of the demonstration
# The signal and negative control files can be used by calling the above
#   command without the writing and reading of the data.
write.table(expressionSignal, 'signal.txt', sep="\t");
write.table(negativeControl, 'negative.control.txt', sep="\t");

#read in those files just created.
data <- mbc.parseFile('signal.txt', 'negative.control.txt');
signal <- data$sig;
negCon <- data$con;
```

---

printMBCBOutput      *MBCB - Model-Based Background Correction for Illumina Beadarray*

---

**Description**

This function is used to neatly output the values created by the other methods in the MBCB package.

**Usage**

```
printMBCBOutput(sig,
  average,
  rmaBool,
  npBool,
  mleBool,
  bayesBool,
  gmleBool,
  avgOutputFile,
  detailOutputFile)
```

**Arguments**

**sig**                    A list corresponding to the background-corrected signal values (as is generated by the `mbcb.correct` function). The list should have elements for all desired background correction methods named 'NP', 'RMA', 'Bayes', and/or 'MLE'.

average	A list containing the average values (as is generated by the <code>mbcb.correct</code> function). The list should have elements for all desired background correction methods named 'NP', 'RMA', 'Bayes', and/or 'MLE'.
npBool	A boolean value representing the desire to compute the non-parametric background correction values.
rmaBool	A boolean value representing the desire to compute the RMA background correction values.
mleBool	A boolean value representing the desire to compute the MLE background correction values.
bayesBool	A boolean value representing the desire to compute the Bayes background correction values.
gmleBool	A boolean value representing the desire to compute the GMLE background correction values.
avgOutputFile	The <i>base</i> file name to which suffixes and a file extension will be appended (i.e. 'C:/output'). These files will store the average values of each background correction method selected.
detailOutputFile	The <i>base</i> file name to which suffixes and a file extension will be appended (i.e. 'C:/output'). These files will store the background corrected intensities

**Value**

This function prints corresponding CSV files based on which background-correction methods were selected.

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**See Also**

[mbcb.main](#)

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