Analysis of Bead Summary Data using beadarray

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Introduction

There are two methods for describing the results of a BeadArray experiment. Firstly, we can use bead-level data whereby the position and intensity of each individual bead on an array is known. The methods available for processing bead level data are discussed in: Dunning, M.J et al, Quality Control and Low-level Statistical Analysis of Illumina Beadarrays, Revstat 4, 1-30 and in a separate vignette of beadarray.

Bead summary data can also be used whereby a summary intensity for each bead type on an array is given. The summarised values for a particular bead type can then be compared between different arrays within an experiment. This is the format of the data output by Illumina's BeadStudio application. The methods described within this document focus exclusivly on dealing with bead summary data. The bead summary data can be data obtained using either the BeadChip (6 or 8 arrays on a slide) or SAM (arrays organised in 96 well plates) technologies.

At present, beadarray is for the analysis of Illumina expression data only. For a package to analyse Illumina SNP data, see beadarraySNP.

1 Citing beadarray

If you use beadarray for the analysis or pre-processing of BeadArray data please cite:

Dunning M, Smith M, Thorne NP, Tavaré, beadarray: An R package to Analyse Illumina BeadArrays, R News, submitted

2 Importing Bead Summary Data

An example data set is included with the beadarray package and can be found as a zipped folder dat directory of the beadarray download. Inside this folder you will find three Excel data files and two text files. The Excel files are the raw data, a sample sheet and a quality control file for the example experiment. These data were obtained as part of a pilot study into BeadArray technology and comprises of 3 Human-6 BeadChips with 6 different samples, I, MC, MD, MT, P and Norm hybridised. MC, MD, MT and P are all tumours whereas Norm is a normal sample and I is a sample provided by Illumina.

2.1 Description of Files

Reading bead summary data into beadarray requires the three files as given for this example experiment and we now describe these in more detail.

• raw_data.csv - This contains the raw, non-normalised bead summary values as output by Bead-Studio and is readable by Excel. Inside the file are several lines of header information followed by a data matrix with some 48,000 rows. Each row is a different gene in the experiment and

the columns give different measurements for the gene. For each array, we record the summarised expression level (AVG_Signal), standard error of the bead replicates (BEADSTDEV), Number of beads used (Avg_NBEADS) and a Detection score which estimates the probability of a gene being detected above the background. Note that whilst this data has not been normalised, it has been subjected to local background correction at the bead level prior to summarising.

- raw_data_sample_sheet Defines the array IDs and samples placed on each array.
- raw_data_qc_info Gives the summarised expression values for each of the controls that Illumina place on arrays and hence extremely useful for diagnostic purposes.

The following code can be used to read the example data into R. Firstly, we have to use the targets.txt file to define the location of the raw data, sample sheet and quality control file. Once this targets information has been read into R we can simply run the function readBeadSummaryData. The default parameters for this function will look for the column headings as described above.

3 The BSData object

BSData is an object of type ExpressionSetIllumina which is an extension of the ExpressionSet class developed by the Biocore team used as a container for high-throughput assays. The data from the the raw_data file has been written to the assayData slot of the object, whereas the phenoData slot contains information from sample_sheet and the QC slot contains the quality control information. For consistency with the definition of other *ExpressionSet* objects, we now refer to the expression values as the *exprs* matrix which can be accessed using exprs and subset in the usual manner. The BeadStDev matrix can be accessed using se.exprs. The rows of exprs are named according to the row names of the original raw_data file.

```
> BSData
Instance of ExpressionSetIllumina
assayData
  Storage mode: list
  Dimensions:
         BeadStDev Detection exprs NoBeads
             47293
                        47293 47293
                                       47293
Features
Samples
                18
                           18
                                 18
                                          18
phenoData
  rowNames: I.1, IC.1, IH.2, ..., Norm.2, P42.2 (18 total)
```

```
varLabels and descriptions:
   Sample_Name: Sample_Name
   Sample_Well: Sample_Well
   Sample_Plate: Sample_Plate
   Sample_Group: Sample_Group
   Pool_ID: Pool_ID
   Sentrix_ID: Sentrix_ID
   Sentrix_Position: Sentrix_Position
featureData
  featureNames: GI_10047089-S, GI_10047091-S, GI_10047093-S, ..., thrB, trpF (47293 total)
  varLabels and descriptions:
Experiment data
 Experimenter name:
 Laboratory:
  Contact information:
 Title:
 URL:
 PMIDs:
 No abstract available.
Annotation [1] "Illumina"
QC Information
Available Slots: Signal StDev Detection
 featureNames: 1475542110_F, 1475542113_E, 1475542114_A, ..., 1475542113_D, 1475542113_F
  sampleNames: Biotin, cy3_high, cy3_low, ..., pm, negative
> exprs(BSData)[1:10, 1:2]
                I.1
                     IC.1
               87.8 131.8
GI_10047089-S
GI_10047091-S 161.8 130.8
GI_10047093-S 481.2 401.4
GI_10047099-S 633.7 483.8
GI_10047103-S 1535.6 1186.5
GI_10047105-S 247.5 210.2
GI_10047121-S 113.0 101.3
GI_10047123-S 453.9 306.8
GI_10047133-A 103.6 114.5
GI_10047133-I 118.0 123.1
> se.exprs(BSData)[1:10, 1:2]
             AVG_Signal.I.1 AVG_Signal.IC.1
GI_10047089-S
                       5.1
                                        9.5
                                        7.9
GI_10047091-S
                       12.0
GI_10047093-S
                       21.7
                                       24.5
                      21.6
                                      20.9
GI_10047099-S
                       42.7
                                       34.5
GI_10047103-S
GI_10047105-S
                      12.7
                                       11.8
GI_10047121-S
                        6.4
                                        8.1
```

GI_10047123-S	14.0	13.1 6.0	
GI_10047133-A	6.8		
GI 10047133-I	5.6	7.2	

> pData(BSData)[, 1:6]

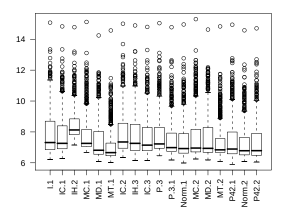
	Sample_Name	Sample_Well	Sample_Plate	Sample_Group	Pool_ID	Sentrix_ID
I.1	NA	NA	NA	IH-1	NA	1475542114
IC.1	NA	NA	NA	IC-1	NA	1475542114
IH.2	NA	NA	NA	IH-2	NA	1475542114
MC.1	NA	NA	NA	MC-1	NA	1475542114
MD.1	NA	NA	NA	MD-1	NA	1475542114
MT.1	NA	NA	NA	MT-1	NA	1475542114
IC.2	NA	NA	NA	IC-2	NA	1475542110
IH.3	NA	NA	NA	IH-3	NA	1475542110
IC.3	NA	NA	NA	IC-3	NA	1475542110
P.3	NA	NA	NA	P-3	NA	1475542110
P.3.1	NA	NA	NA	P-3	NA	1475542110
Norm.1	NA	NA	NA	Norm-1	NA	1475542110
MC.2	NA	NA	NA	MC-2	NA	1475542113
MD.2	NA	NA	NA	MD-2	NA	1475542113
MT.2	NA	NA	NA	MT-2	NA	1475542113
P42.1	NA	NA	NA	P-1	NA	1475542113
Norm.2	NA	NA	NA	Norm-2	NA	1475542113
P42.2	NA	NA	NA	P-2	NA	1475542113

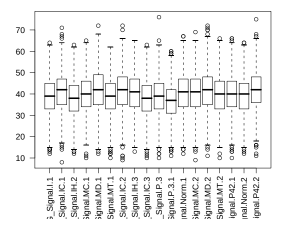
Boxplots of expression may be useful for quality control. Below we show the code to produce boxplots of the log2 intensities of each array in the experiment. Recall that there are 6 arrays per BeadChip and that differences between chips hybridisations on different days may be expected. In this example the differences in intensity between arrays on the same chip and different chips do not seem too large. However, we can see that the first BeadChip seems to be more variable than the others and in particular the third array on the first BeadChip could be an outlier.

Boxplots of the other slots in BSData can be easily plotted. 1

```
> par(mfrow = c(1, 2))
> boxplot(log2(exprs(BSData)[1:1000, ]), las = 2)
> boxplot(NoBeads(BSData)[1:1000, ], las = 2)
```

¹We have restricted the number of points plotted in order to keep the size of this vignettee small.

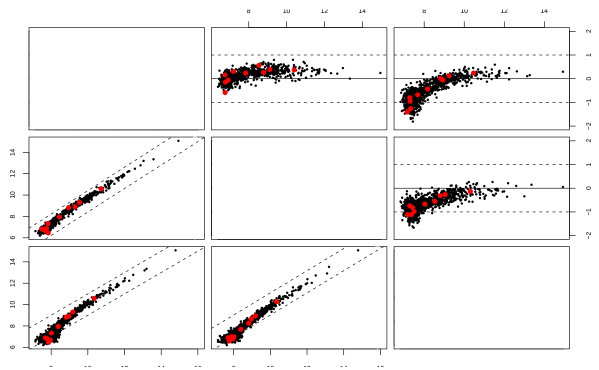




4 Normalisation and Quality Control

In the expression boxplots we notice that there are differences in expression level across a chip and between chips. Therefore we might want to normalise the arrays in the experiment comparable. We also see the the 3rd array has significantly different intensity. The sample on this array is replicated three times on the first chip in the experiment, so comparing the MA and XY plots for the sample can be informative about the variability of this sample.

Particular genes of interest may be highlighted on the MA and XY plots by using the <code>genesToLabel</code> argument which should match up with the row names in <code>BSData</code>. The <code>labelCol</code> argument can be used to specify a colour for each gene. For simplicity sake we simply highlight the first ten genes in the expression matrix, a possible application might be to highlight control genes on the plot or particular genes of interest.



In the top right corner we see the MA plots for all pairwise comparisons involving the 3 arrays. On an MA plot, for each gene we plot the average of the expression levels on the two arrays on the x axis and the difference in the measurements on the y axis. For replicate arrays we would expect all genes to be unchanged between the two samples and hence most points on the plot to lie along the line y=0. In the lower left corner of the MAXY plot we see the XY plot and for replicate arrays we would expect to see most points along the diagonal y = x. From this MAXY plot it is obvious that the third array is significantly different to the other replicates and should probably be discarded.

Both XY and MA plots for a particular comparison of arrays are available separately using plotXY and plotMA

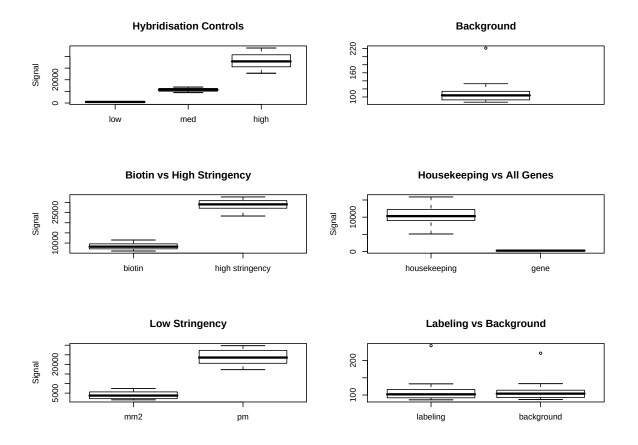
The quality control information which is read in by readBeadSummaryData can be plotted to provide useful diagnostic information. To retrieve this quality control data we can use the QCInfo function. Alternatively, quality control information can be read using readQC.

The QC object contains Signal, StDev and Detection matrices with each row in the matrix being a different array and each column a different control type. An overview of QC can be plotted using plotQC.

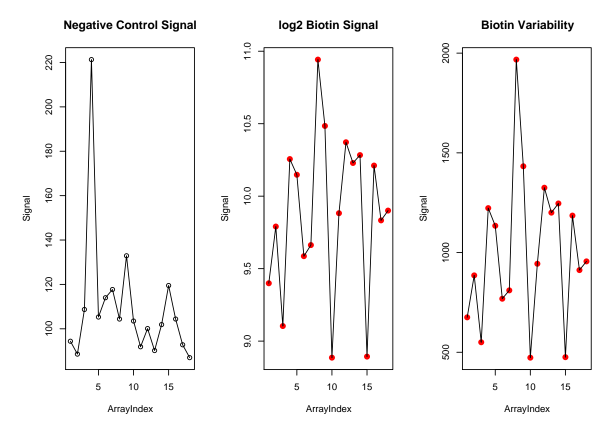
```
> QC = QCInfo(BSData)
> QC$Signal[1:3, ]
```

```
Biotin cy3_high cy3_low cy3_med gene
                                                               house labeling
                                816.6 11178.2 205.8 29498.3
                                                              7914.2
                                                                          92.9
1475542110_F
              7551.0
                      32436.0
1475542113_E 6137.2
                      28081.0
                                739.4 9158.1 176.6 23302.4
                                                               6680.7
                                                                          86.1
1475542114_A 10255.0
                      41451.7
                               1040.9 13176.7 320.3 30390.5 15902.3
                                                                         106.0
                         pm negative
                 mm
1475542110_F 3584.5 21807.1
                                 94.4
1475542113_E 1516.5 18619.5
                                88.6
1475542114_A 5738.7 27314.2
                                108.7
```

> plotQC(BSData)



The singleQCPlot function allows a particular control type to be plotted across all samples. The type argument must match one of the column names of QC\$Signal and the what argument selects which of the Signal, StDev and Detection slots to plot. Additional plotting arguments such as a title for the plot, plotting character etc can also be passed to the function. We can also choose to plot on the log₂ scale.

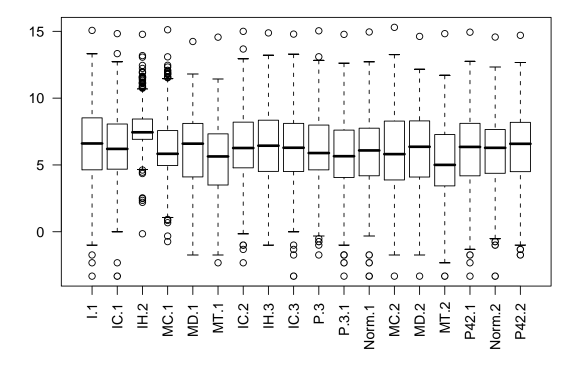


Illumina also use this quality control information to normalise bead summary data. In a procedure known as background normalisation, the averaged values of all negative controls on a particular array are subtracted from the summarised expression of each gene. This normalisation can be repeated by the function backgroundNormalise. The intended effect of this normalisation is to remove the effects of non-specific binding from the expression values. This effect is more noticable for genes with low expression level and hence can produce negative values.

```
> BSData.bgnorm = backgroundNormalise(BSData)
```

- [1] 65.9 45311.7
- > range(exprs(BSData.bgnorm)[, 1])
- [1] -142.7 45221.5
- > boxplot(log2(exprs(BSData.bgnorm)[1:1000,]), las = 2)

> range(exprs(BSData)[, 1])



It is possible to use the normalisation methods available in the affy such as quantile, qspline or others. The method of rank invariant normalisation recommended by Illumina may also be applied once a suitable target distribution has been defined. In the following example we define this to be the mean of each row before using the normalize.invariantset to find a set of invariant genes and define a normalising curve using this set and the target distribution.

```
> library(affy)
> BSData.quantile = assayDataElementReplace(BSData, "exprs", normalize.quantiles(as.matrix(exprs(BSData))))
> BSData.qspline = assayDataElementReplace(BSData, "exprs", normalize.qspline(as.matrix(exprs(BSData))))
> T = apply(exprs(BSData.bgnorm), 1, mean)
> BSData.rankinv = assayDataElementReplace(BSData.bgnorm, "exprs",
+ rankInvariantNormalise(exprs(BSData.bgnorm), T))
```

5 Differential Expression

Research into the best method for detecting differential expression for BeadArray data is still work in progress. In the meantime, users are able to use the lmFit and eBayes functions from limma on the matrix exprs(BSdata) with a log₂ transformation applied.

The following code shows how to set up a design matrix for the example experiment combining the I, MC, MD, MT, P and Normal samples together. We then define contrasts comparing the I samples to the P samples and I to Normal and perform an empirical bayes shrinkage. In this particular experiment, the I and P samples are completely different so we would expect to see plenty of differentially expressed genes.

For more information about 1mFit and eBayes please see the comprehensive limma documentation.

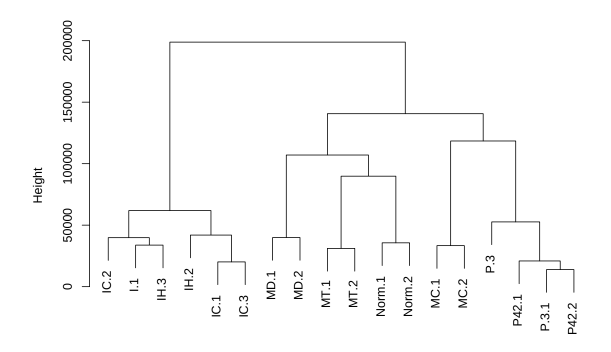
The algorithm for the Illumina method is implemented in the function DiffScore although it not completely accurate at present. To compare array 1 in the experiment to array 10 (ie comparing an I sample to a P) we would use the following which seems to pick a lot of genes with significantly high expression on array 10 than array 1.

```
> df = DiffScore(BSData, QC, cond = 10, ref = 1)
> o = order(df, decreasing = TRUE)[1:50]
> exprs(BSData)[o, 10]/exprs(BSData)[o, 1]
 [1] 81.903596 58.697624 26.983822 26.503597 18.531785 27.010807 30.735270
 [8] 13.053541 22.707432 20.873933 13.656178 17.296035 10.780707 12.992251
[15] 12.161406 9.797336 14.039683 10.480079 14.698404 13.677614 10.798242
[22] 10.442509 11.538138 6.694895 6.355486 6.817261
                                                       8.692986
[29] 10.223131 5.777421 4.528634 5.195435 10.384192
                                                       7.898985
                                                                 4.903955
[36]
               4.524027 5.929239 6.135308 4.707682 6.059928
     5.885077
                                                                 5.306415
[43]
     4.732153 6.848549 9.298222 4.262486 3.899440 3.870005
[50] 3.677042
```

6 Further Analysis

The clustering functionality available in BeadStudio can be easily performed through R using the hlcust once a distance matrix has been defined. In this example we see that the clusters correspond well to the different sample types. The heatmap function could also be used in a similar manner and principal components analysis is possible using princomp.

```
> d = dist(t(exprs(BSData)))
> plclust(hclust(d), labels = rownames(pData(BSData)))
```



d hclust (*, "complete")