EMBO03 Lab4: Introduction to Bioconductor affy Package

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In this lab, we demonstrate the main functions in the affy package for pre-processing Affymetrix microarray data. For a more detailed introduction, consult the package vignettes which can be listed by the command openVignette("affy"). A demo can also be accessed by demo(affy). A number of sample datasets are available in the package; to list these, type data(package="affy"). To load the package

> library(affy)

The function ReadAffy is available for reading CEL files. However, in this lab we will work mainly with the Dilution dataset, which is included in the package. For a description of Dilution, type ? Dilution. To load this dataset

> data(Dilution)

One of the main classes in affy is the AffyBatch class. For details on this class consult the help file, ? AffyBatch; methods for manipulating instances of this class are also described in the help file. Other classes include ProbeSet (PM and MM intensities for individual probe sets), Cdf (information contained in a CDF file), and Cel (single array cel intensity data). The object Dilution is an instance of the class AffyBatch. Try the following commands to obtain information on this object

> class(Dilution)

```
[1] "AffyBatch"
```

```
> slotNames(Dilution)
```

[1]	"cdfName"	"nrow"	"ncol"	"exprs"	"se.exprs"
[6]	"phenoData"	"description"	"annotation"	"notes"	

> Dilution

```
AffyBatch object
size of arrays=640x640 features (12805 kb)
cdf=HG_U95Av2 (12625 affyids)
number of samples=4
number of genes=12625
annotation=hgu95av2
notes=
```

> annotation(Dilution)

[1] "hgu95av2"

For a description of the target samples hybridized to the arrays

> phenoData(Dilution)

phenoData object with 3 variables and 4 cases varLabels

liver: amount of liver RNA hybridized to array in micrograms sn19: amount of central nervous system RNA hybridized to array in micro scanner: ID number of scanner used

> pData(Dilution)

	liver	sn19	scanner
20A	20	0	1
20B	20	0	2
10A	10	0	1
10B	10	0	2

The exprs slot contains a matrix with columns corresponding to arrays and rows to individual probes on the array. To obtain the matrix of intensities for all four arrays

```
> e <- exprs(Dilution)
> nrow(Dilution) * ncol(Dilution)
```

[1] 409600

> dim(e)

```
[1] 409600 4
```

You can access probe-level PM and MM intensities using

```
> PM <- pm(Dilution)
> dim(PM)
```

```
[1] 201800
                4
> PM[1:5, ]
                 20B
           20A
                        10A
                              10B
1000_at1 468.8 282.3 433.0 198.0
1000_at2 430.0 265.0 308.5 192.8
1000_at3 182.3 115.0 138.0 86.3
1000_at4 930.0 588.0 752.8 392.5
1000_at5 171.0 128.0 152.3 97.8
  To get the probe set names (Affy IDs)
> gnames <- geneNames(Dilution)</pre>
> length(gnames)
[1] 12625
> gnames[1:5]
[1] "1000_at"
                           "1002_f_at" "1003_s_at" "1004_at"
                "1001_at"
> nrow(e)/length(gnames)
```

[1] 32.44356

As with other microarray objects in Bioconductor packages, you can use subsetting commands for AffyBatch objects

```
> dil1 <- Dilution[1]
> class(dil1)
[1] "AffyBatch"
> dil1
AffyBatch object
size of arrays=640x640 features (3204 kb)
cdf=HG_U95Av2 (12625 affyids)
number of samples=1
number of genes=12625
annotation=hgu95av2
notes=
> cel1 <- Dilution[[1]]
> class(cel1)
```

```
[1] "Cel"
> cel1
Cel object
name=20A
cdfName=HG_U95Av2
intensity=640 x 640 (3200 kb)
masked= 0 %
outliers= 0 %
history=
```

One of the main functions for reading in Affymetrix data is ReadAffy. It reads in data from CEL and CDF files and creates objects of class AffyBatch. Using ReadAffy(widget=TRUE) provides widgets for interactive data input.

To produce a spatial image of probe log intensities and probe raw intensities

```
> image(Dilution[1])
```



20A

> image(cel1)

To produce boxplots of probe log intensities

> boxplot(Dilution, col = c(2, 2, 3, 3))



Small part of dilution study

To produce density plots of probe log intensities

> hist(Dilution, type = "l", col = c(2, 2, 3, 3), lty = rep(1:2, + 2), lwd = 3)



The boxplots and density plots show that the Dilution data needs normalization. As described in the dataset help file and in the phenoData slot (pData(Dilution)), two concentrations of mRNA were used and, for each concentration, two scanners were used. From the plots, we note that scanner effects seem stronger than concentration effects (different colors). Arrays that should be the same are different; arrays that should be different are similar. Because different mRNA concentrations were used, we perform probe-level normalization within concentration groups.

```
> Dil20 <- normalize(Dilution[1:2])
> Dil10 <- normalize(Dilution[3:4])
> Dil10 <- normalize(Dilution[3:4])</pre>
```

```
> normDil <- merge(Dil20, Dil10)</pre>
```

Notice how the boxplot now looks better.

```
> boxplot(normDil, col = c(2, 2, 3, 3))
```

Small part of dilution study



The affy package provides implementations for a number of methods for background correction, probe-level normalization (e.g., quantile, curve-fitting (Bolstad et al., 2002)), and computation of expression measures (e.g., MAS 4.0, MAS 5.0, MBEI (Li & Wong, 2001), RMA (Irizarry et al., 2003)). To list available methods for AffyBatch objects

```
> bgcorrect.methods
```

```
[1] "mas" "none" "rma"
> normalize.AffyBatch.methods
[1] "constant" "contrasts" "invariantset" "loess"
[5] "qspline" "quantiles" "quantiles.robust"
> pmcorrect.methods
[1] "mas" "pmonly" "subtractmm"
> express.summary.stat.methods
```

	[1]	"avgdiff"	"liwong"	"mas"	"medianpolish"	"playerout"
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The main normalization function is expresso. You can select pre-processing methods interactively using widgets by typing expresso(Dilution, widget=TRUE). The function operates on objects of class AffyBatch and returns objects of class exprSet. rma provides a more efficient implementation of Robust Multi-array Average (RMA). We don't normalize because we already did above.

```
> rmaDil <- rma(normDil, normalize = FALSE)
> class(rmaDil)
```

Data packages for CDF information can be download from www.bioconductor.org. These packages contain environment objects which provide mappings between AffyIDs and matrices of probe locations, with rows corresponding to probe-pairs and columns to PM and MM cels. CDF environments for HGU95Av2 and HGU133A chips are already in the package. For information on the environment object ? hgu95av2cdf

```
> annotation(Dilution)
[1] "hgu95av2"
> data(hgu95av2cdf)
> pnames <- ls(env = hgu95av2cdf)</pre>
> length(gnames)
[1] 12625
> gnames[1:5]
[1] "1000_at"
                "1001_at"
                          "1002_f_at" "1003_s_at" "1004_at"
> get(gnames[1], env = hgu95av2cdf)
          pm
                 mm
 [1,] 358160 358800
 [2,] 118945 119585
 [3,] 323731 324371
 [4,] 223978 224618
 [5,] 313420 314060
 [6,] 349209 349849
 [7,] 199525 200165
 [8,] 213669 214309
 [9,] 236739 237379
[10,] 298099 298739
[11,] 282744 283384
```

[12,] 281443 282083
[13,] 349198 349838
[14,] 297953 298593
[15,] 317054 317694
[16,] 404069 404709

You can also use the indexProbes, pmindex, and mmindex functions to get information on probe location

```
> plocs <- indexProbes(Dilution, which = "both")</pre>
> plocs[[1]]
 [1] 358160 118945 323731 223978 313420 349209 199525 213669 236739 298099
[11] 282744 281443 349198 297953 317054 404069 358800 119585 324371 224618
[21] 314060 349849 200165 214309 237379 298739 283384 282083 349838 298593
[31] 317694 404709
> pmindex(Dilution, genenames = gnames[1], xy = TRUE)
$"1000_at"
        х
            У
 [1,] 400 560
 [2,] 545 186
 [3,] 531 506
 [4,] 618 350
 [5,] 460 490
 [6,] 409 546
 [7,] 485 312
 [8,] 549 334
 [9,] 579 370
[10,] 499 466
[11,] 504 442
[12,] 483 440
[13,] 398 546
[14,] 353 466
[15,] 254 496
[16,] 229 632
> pmindex(Dilution, genenames = gnames[1])
$"1000_at"
 [1] 358160 118945 323731 223978 313420 349209 199525 213669 236739 298099
```

[11] 282744 281443 349198 297953 317054 404069

Having access to PM and MM data can be useful. Let's look at a plot of PM vs. $\rm MM$

```
> plot(mm(Dilution[1]), pm(Dilution[1]), pch = ".", log = "xy")
> abline(0, 1, col = "red")
```



mm(Dilution[1])