GS01 0163
Analysis of Microarray Data

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Lecture 8: Normalization and Affymetrix

- What is Normalization?
- What Methods have People Suggested?
- How Can These Methods be Tested?
- What do we Recommend?
- Revisiting Normalization in BioConductor
- R manipulations of AffyBatch
What is Normalization?

Broad question: How do we compare results across chips?

Focused goal: Getting numbers (quantifications) from one chip to mean the same as numbers from another chip.
Why is Normalization an Issue?

Amount of RNA

Efficiencies of RNA extraction, reverse transcription, labeling, photodetection

PCR yield

DNA quality
Why is Normalization an Issue?

Variation that is obscuring as opposed to interesting.
What Has Been Tried?

Housekeeping genes – start with a set of genes whose expression you believe shouldn’t change, and scale the other expression values accordingly.

Spike-ins – introduce a set of markers whose relative intensities you can control, and use these to calibrate the remaining intensities.

Simple scaling – multiply all of the intensities from one chip so that a summary of the result matches the summary value from another chip or some standard.
Some Difficulties

Housekeeping genes – what are they? Are there any genes whose expression does not change across a wide variety of conditions? Do those conditions include cancer (broad distortion)?

Spike-ins – How do we regulate the amount of the spike-in relative to the amount of the material of interest?

Simple scaling – Does a log scale MA plot look flat? What scaling factor do we use? Median? Total amount of RNA present? Some other quantile?
Non-linear normalization

N18__normal.CEL vs N01__normal.CEL

Median: 0.22
IQR: 0.598
An Alternative?

Combining Spiking and scaling?

Use spike-ins covering a very broad dynamic range, and use this to try to define “linearity” of expression, after which you scale.
Housekeeping: Can this be Inferred?

What is a housekeeping gene?

Typical assumption – something that is vital to the stable functioning of the cell, present at constant levels. In most suggested cases, this level has been high.

Slightly different definition – a gene that retains roughly the same rank in a sorted list of expression values.

Choose a subset with “invariant ranks” and use these to determine our mapping (Schadt et al.)
What is Our Baseline?

When we are assessing invariance, we have to define “with respect to what?”

For the most part, this has meant choosing a single chip as a canonical “reference” or “baseline”.

Typically, this chip is taken to be a “middle” chip by some metric, such as the chip having the median “median intensity”.

This is the default in dChip.
Invariance and Nonlinearity

Using an invariant set effectively applies a whole bunch of scaling factors, with the factor changing depending on the intensity (scaling at high levels is different than scaling at low levels).

For the most part, this serves to map the quantiles of one set of intensities over to another.

Similar to producing an MA-plot, fitting a smooth curve (e.g. loess) to the center of the data, and subtracting the curve off.
An Implicit Assumption

Is matching the 90th percentile reasonable if one sample causes no gene expression whatsoever, and a second produces expression in half of the genes?

The assumption that mapping using quantiles or scaling is reasonable is based on the assumption that “most genes don’t change”, and quantiles use this more extensively than scaling.

If this underlying assumption is doubtful, then using the above methods is shaky.
Cases Where Most Genes Do Change

Fed/Starved experiments

Heat shock experiments

Experiments comparing different organs
How Can We Test Normalization Methods?

Most of the following is based on a paper by Bolstad, Irizarry, Astrand and Speed (Bioinformatics, 2003, p.185-193)

When can we know what the true answer should be?

If we’re looking at the same stuff, or stuff that differs only in a way that we view as irrelevant to the biology, then things should look the same after normalization.

Find some good datasets that work like this.
Affymetrix Data

The Affymetrix Latin Square Experiment (Hu95Av2, repeated on Hu133A)

roughly 42 chips which have the same stuff printed modulo changes in spike-ins.
Gene Logic Dilution Data

75 Hu95Av2 arrays, two tissue sources: liver and central nervous system.

30 chips each for each tissue alone:

• in 5 blocks of size 6, with each block at a different dilution level.

Remaining 15 chips are mixtures of liver and CNS:

• in 3 blocks of 5, with ratios of 25:75, 50:50, and 75:25.

4 of the liver arrays are included with the BioConductor affydata package (Dilution).
A Gene Logic Spike In Dataset

98 Hu95Av1 arrays, with 11 cRNA spike-ins. For 26 of these chips, with a common AML source, the level was nominally the same for all of the spike-ins, but this level was altered from chip to chip (i.e., chip 1 had all 11 at 0, chip 2 had all 11 at 0.5, chip 3 at 0.75, etc.) Remaining chips were set up in two Latin Square experiments with 3 replicates at each level.

With the first 26, nothing should change except the spike-ins.
What was Used?

5 of the liver arrays from the Dilution experiment (all at the same dilution level)

Should normalization correct for dilution?

all 26 arrays from the spike-in experiment where the spike-in levels were nominally the same.
Aside: Why did Gene Logic Do This?

Gene Logic is the biggest single consumer of Affy chips, and they have assembled very large databases of expression profiles, which they market.

They are trying to improve the analysis (which helps them) and to advertise their own approaches (again).

The datasets can be requested (they’ll send you a cd).
Given Data, What Tests Should we Use?

Baseline methods:

- Scaling a la Affy, equating 96% trimmed means and choosing a baseline chip (median of median). (Affy’s algorithm performs this scaling on the summary measures, but probe levels were used here).

- Nonlinear invariant rank approach, a la Schadt et al., again using a baseline chip (the same one).

Some “Complete Data Methods” that they introduce.
Complete Data: Eliminating Baseline

Cyclic Loess

Contrasts

Quantiles

All of these work by treating the chips in a symmetric fashion, and all of them are implemented as normalization methods in the BioConductor affy package.
Cyclic Loess Normalization

Start with MA plots, fit a loess smooth for each pair of chips.

Let $M_k = \log_2(x_{ki}/x_{kj})$ for arrays $i, j$, and let $\hat{M}_k$ denote the fitted loess curve for this pair of chips. Then the adjusted value should be $M'_k = M_k - \hat{M}_k$.

How much should we adjust each of the chips? Use a symmetric approach

$$x'_{ki} = 2^{A_k + 0.5 \times M'_k}, \quad x'_{kj} = 2^{A_k - 0.5 \times M'_k}$$

Repeat for all pairs, then refit and repeat. (2-3 iterations required.)

This is Slow (at least $O(n^2)$ for $n$ chips).
Contrast Based Normalization

Start with vectors of log intensities from each chip. Use a rotation matrix (a matrix of orthogonal *contrasts*) to change the basis. The first column of the converted matrix is the average of all of the log intensities. For the remaining $n - 1$ columns ($n$ arrays), fit loess curves, flatten out, and reverse the transformation.

Central focus is the average on the log scale (the geometric mean).

Faster than cyclic loess, but still slow.
Matrixes before rotation
Matrixes after rotation
Quantile Normalization

Assume that the distributions of probe intensities should be completely the same across chips.

Start with $n$ arrays, and $p$ probes, and form a $p \times n$ matrix $X$.

Sort the columns of $X$, so that the entries in a given row correspond to a fixed quantile.

Replace all entries in that row with their mean value, and undo the sort.

This procedure, sorting and averaging, is comparatively fast.
Quantile Argument - Why the Mean?

Why did they choose the mean? If we work in 2d, then perfect quantile agreement produces a straight line.

If we plot $n$-vectors of quantiles, then we want to see a straight line in $n$-space along the main diagonal.

Projecting the observed $n$-vector onto this central axis suggests using the mean value.
Our Quibble: Scale

The argument suggesting the mean does not address the issue of what scale the observations are on, and works equally well for both the raw and log scales, but the mean values differ.

We prefer the median, which is much more scale-invariant, but for the most part there is little practical difference.
Does Baseline Matter?

Actually, yes.

from Astrand (J. Comp. Biol., 2003, p.95-102)

Starting with a pair of arrays (Mu11K), they normalized in dChip using array A as baseline, and again using array B as baseline.

Of the predicted log ratios, roughly 18% of the ratios (1603/8799) differed by more than 10%.
A Related View...

Of the lists produced using dChip’s filters, there were 469 genes one way, and 298 the other.

The overlap was 265.

We have seen similar results with more chips.
How Do They Define Success?

We want the results for a given probeset to be the same from one chip to the next.

For each method, compute expression values, (put them on the log scale ?), and compute the variance and the average (this is akin to what is required for a multivariate MA plot).

These are our method summaries.
How Do They Define Success?

For each pair of methods, produce a plot with the average of the log means on the x-axis and the log ratio of the variances on the y-axis.

Fit a smooth curve, and look for cases where one method’s variance is much smaller than the other’s.
What Preprocessing Did They Employ?

To make sure that differences in the summary method employed do not drive the results, pick one method and use it throughout.

They use RMA with the default BG correction throughout.
Some Results

In general, pairwise MA plots did show some curvature before normalization, suggesting that a nonlinear fit might help.

All of the methods that they used reduced the variance of the numbers substantially from using just the raw numbers without normalization.

The complete data methods also improve on the baseline chip methods, and they improve more on simple constant scaling than they do on nonlinear fits such as the invariant rank set.
More Results

With respect to the three “complete data” methods, the differences are fairly small, with the quantile method looking very slightly better.

As the quantile method is definitely the fastest, this is the one that they recommend.
Do We Agree?

Of course!

In this case, the assumptions underlying this type of approach are met, and most genes should not change.

But this assumption is not always met, and in cases where we are suspicious of this we try to use something that is less dependent on a large number of assumptions.

This can be simple scaling, or scaling for both mean and spread.
What Do We Do?

In general, the first thing we do is check the context of the experiment, to see if we are persuaded that most genes might be the same.

Next, we check the pairwise MA plots. If we don’t see substantial nonlinearity, then we don’t have to use quantiles so we don’t.

If we do see nonlinearity, we check images of the CEL files to look for artifacts, and if we see nothing we’ll go with quantiles.
A Bioconductor Adventure...

Our goal – to reproduce the study of Bolstad et al. (2003) using the data supplied with BioConductor.

First, pull in the Affy functions and get the data

> library(affy);
> library(affydata);
> data(Dilution);
What steps are we trying to follow?

Starting with an AffyBatch object, presumably assembled straight from CEL files, we want to test the effects of different normalization methods on the stability of probeset measurements of the same stuff.

The steps:

Background correction
Normalization
PM correction
Summary Quantification

Monitor as we go!
boxplot(Dilution); # shows log intensities!
dev.copy(png, file="boxplot1.png", col=2:5);
dev.off();
What about the densities? (View 2)

```r
hist(Dilution, lty=1, col=2:5, lwd=3);
dev.copy(png, file="hist1.png");
dev.off();
```
and the MA plots?

```r
par(mfrow=c(2,2));
MAplot(Dilution);
par(mfrow=c(1,1));
```
Look at all pairs?

```r
mva.pairs(Dilution);
```

Error in \( \log(x, \text{base}) \) : Non-numeric argument to mathematical function

> `help(mva.pairs)`

want to feed this function a matrix, with columns corresponding to arrays. Where are these numbers?
I can never remember...

Objects have slots!

> slotNames(Dilution)
[1] "cdfName" "nrow" "ncol"
[4] "assayData" "phenoData" "featureData"
[7] "experimentData" "annotation" ".__classVersion"

We can extract the numbers we want with `exprs`.

> length(exprs(Dilution))
[1] 1638400
> dim(exprs(Dilution))
[1] 409600 4
What’s in the Slots?

Dilution@cdfName
[1] "HG_U95Av2"

> Dilution@nrow
[1] 640

> Dilution@ncol
[1] 640

> Dilution@phenoData
sampleNames: 20A, 20B, 10A, 10B

varLabels and varMetadata:
liver: amount of liver RNA hybridized to array
sn19: amount of central nervous system RNA hybridized to array
scanner: ID number of scanner used

> Dilution@experimentData
Experiment data
Experimenter name: Gene Logic
Laboratory: Gene Logic
Contact information: 708 Quince Orchard Road
Gaithersburg, MD 20878
Telephone: 1.301.987.1700
Toll Free: 1.800.GENELOGIC (US and Canada)
Facsimile: 1.301.987.1701

Title: Small part of dilution study
URL: http://qolotus02.genelogic.com/datasets.nsf/
PMIDs:
Abstract: A 68 word abstract is available. Use 'abstract' method.
Other:
> Dilution@annotation
[1] "hgu95av2"
> Dilution@.__classVersion__
   R     Biobase     eSet     AffyBatch
   "2.5.0" "1.13.22" "1.1.0"  "1.2.0"

Still haven’t touched assayData or featureData...
What’s in the Slots? (pt. 2)

> Dilution@featureData
An object of class "Annotated Data Frame"
  featureNames: 1, 2, ..., 409600 (409600 total)
  varLabels and varMetadata: none
> class(Dilution@featureData)
[1] "AnnotatedDataFrame"
attr(,"package")
[1] "Biobase"
> slotNames(Dilution@featureData)
[1] "varMetadata" "data" "dimLabels" ".__classVersion__"
> dim(Dilution@featureData@data)
[1] 409600 0
> rownames(Dilution@featureData@data)[10]
[1] "10"
What’s in the Slots? (pt. 3)

> Dilution@assayData
$exprs

20A 20B 10A 10B
1 149.0 112.0 129.0 60.0
...
24999 417.8 305.8 358.0 212.5
[ reached getOption("max.print") -- omitted 384601 rows ]

> class(Dilution@assayData)
[1] "list"

> length(Dilution@assayData)
[1] 1

> names(Dilution@assayData)
[1] "exprs"

> class(Dilution@assayData[[1]])
[1] "matrix"

> dim(Dilution@assayData[[1]])
[1] 409600 4
mva.pairs(exprs(Dilution));
Spatial Plots?

image(Dilution[,1], transfo=log2);

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Ratios of Spatial Plots?

```r
image(matrix(exprs(Dilution[,1]),
            nrow=nrow(Dilution),
            ncol=ncol(Dilution)),
    transfo=log2);
```

Parameter “transfo” can’t be set in high-level plot() function.

```r
image(log2(matrix(
            exprs(Dilution[,1]),...
```
Spatial Plot 1

```r
dilutionPlot <- image(log2(matrix(exprs(Dilution[,1]),..)),
  main=sampleNames(Dilution[,1]));
```
Ratio Plot 1 (problem: fake geometry)

```r
dilution.plot <- image(log2(matrix(exprs(Dilution[,1])/
                                exprs(Dilution[,2]),..)),
               main=paste(sampleNames(Dilution[,1]),...));
```

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Ok, start processing. BG first

Dilution.bg <- bg.correct.rma(Dilution);

Did this change things?

hist(Dilution.bg, lty=1, col=2:5, lwd=3)

Let's also try it a different way to make sure...

plotDensity(log2(exprs(Dilution.bg)),
           lty=1, col=2:5, lwd=3)
Picture 1 After BG

hist(Dilution.bg, lty=1, col=2:5, lwd=3)
plotDensity(log2(exprs(Dilution.bg)), lty=1, col=2:5, lwd=3)
plotDensity(log2(pm(Dilution.bg)),
lty=1,col=2:5,lwd=3)
Is Background a Big Deal?

Dilution.bg <- bg.correct.mas(Dilution);
hist(Dilution.bg, lty=1, col=2:5, lwd=3);
title(main="Dilution data, MAS background");
and now we normalize!

This is where the differences come in. We can invoke

normalize.AffyBatch.constant
normalize.AffyBatch.contrasts
normalize.AffyBatch.invariantset
normalize.AffyBatch.quantiles

or, of course, we can have expresso
Expresso, no normalization

eset0 <- expresso(Dilution,
   bgcorrect.method="rma",
   normalize=FALSE,
   pmcorrect.method="pmonly",
   summary.method="medianpolish");

Now at this point, \texttt{eset0} is an ExpressionSet object; the dimensions of the matrix extracted by \texttt{exprs} have changed as we have shifted from features (probes) to probesets.
What Does an ExpressionSet Have?

```r
> slotNames(eset0)
[1] "assayData"  "phenoData"  "featureData"
[4] "experimentData"  "annotation"
[6] ".__classVersion__"
> rownames(eset0@featureData@data)[10]
[1] "1009_at"
> eset0@assayData
<environment: 0x1cb8f904>
> ls(eset0@assayData)
[1] "exprs"  "se.exprs"
> dim(get("exprs", eset0@assayData))
[1] 12625 4
```
Checking the Environment

> myEnv <- new("environment")
> frogs <- rnorm(5)
> assign("frogs", frogs, envir=myEnv)
> ls(myEnv)
[1] "frogs"

Environments are useful things. Basically, they’re R’s answer to “pass by reference” instead of “pass by value”.

What do we want?

The mean and variance of the probeset measurements gene by gene, to describe the behavior of this normalization method.

```r
> dim(exprs(eset0))
[1] 12625 4
> eset0.mu <- apply(exprs(eset0),1,"mean");
> eset0.var <- apply(exprs(eset0),1,"var");
```

Now we want another method to compare to.
**Constant normalization: choosing baseline**

find the “middle behavior” chip

> apply(exprs(Dilution),2,"median");
20A 20B 10A 10B
188 127 149 94

eset1 <- expreso(Dilution,
   bgcorrect.method = "rma",
   normalize.method = "constant",
   normalize.param = list(refindex=3),
   pmcorrect.method = "pmonly",
   summary.method = "medianpolish");

> eset1.mu <- apply(exprs(eset1),1,"mean");
> eset1.var <- apply(exprs(eset1),1,"var");
So, how do we compute MA plots here?

Normally, we are plotting the results from one chip against that from another. Here, we are working with two sets of results from the same chips, just using different methods for quantification.

\[
A1 \leftarrow \frac{(\text{eset0} \cdot \text{mu} + \text{eset1} \cdot \text{mu})}{2}; \\
M1 \leftarrow \frac{(\text{eset0} \cdot \text{mu} - \text{eset1} \cdot \text{mu})}{2}; \quad \# \text{ not quite.} \\
M2 \leftarrow \frac{\text{eset0} \cdot \text{var} / \text{eset1} \cdot \text{var}}{2}; \quad \# \text{ still not quite.} \\
M3 \leftarrow \log_2(\frac{\text{eset0} \cdot \text{var}}{\text{eset1} \cdot \text{var}}); \\
\]

This may still be driven by outliers. Tweak.

\[
d0 \leftarrow 0.0001; \\
M4 \leftarrow \log_2\left(\frac{(\text{eset0} \cdot \text{var} + d0)}{(\text{eset1} \cdot \text{var} + d0)}\right); \\
\]
Dilution: “none” against “scaling”

Here, 12615 times out of 12625, constant scaling gives lower variability. Mean log diff: 4.65
Dilution: “scaling” against “quantiles”

Here, 9477 times out of 12625, quantile scaling gives lower variability. Mean log diff: 0.98
What didn’t they do?

Our comparison of normalization methods here focused on reducing variability, and it assumed that a particular type of background correction (rma) and summarization (median polish) had been employed.

But we saw that different background correction methods led to different shapes in the distributions of probe intensities. If we use “mas” as the background subtraction method, are the differences between the normalization methods still as stark?
Dilution: “none” against “scaling”, MAS BG

Here, 12600 times out of 12625, constant scaling gives lower variability. Mean log diff: 5.40
Dilution: “scaling” vs “quantiles”, MAS BG

Here, 7937 times out of 12625, quantile scaling gives lower variability. Mean log diff: 0.265
What Method Works Best

- AffyComp (http://affycomp.jhsph.edu/) is a web-based tool for comparing competing expression measures.

- Competitors return expression-level data, and they assess how well their method performs.