### BIOS 7659 Journal Club:

A comparison of normalization methods for high density oligonucleotide array data based on variance and bias. (Bolstad et al., 2003)

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#### Introduction

- ► The goal of normalization is to separate the interesting biological variation from the variation that is a result of sample preparation, array production and processing, etc.
- Affymetrix proposes scaling the arrays so that each one has the same mean expression (based on a summary measure).
  - ► This does not work well when there are non-linear relationships between arrays.

### **Alternatives**

- Other approaches, such as non-linear smooth curves or transforming data to standardize the distribution of intensities across arrays, rely on picking a "baseline" array.
- Bolstad et al. compare three different approaches, all of which combine data from every single array (complete data).

## Cyclic loess

- ▶ Basically an extension of the M vs. A plots discussed in class, but applied to pairwise combinations of Affymetrix arrays.
- ▶ M is the difference in log expression values and A is the average (a Bland-Altman plot).

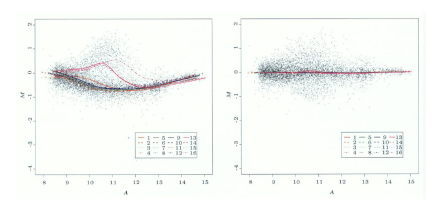


Figure 1: Dudoit et al., 2002

# Cyclic loess

- 1. Take two arrays i and j, each with probes k = 1, ..., p.
- 2. Create an MA plot for these two arrays, and fit a loess curve through these data:

$$M_k = log_2(\frac{x_{ki}}{x_{kj}}), A_k = \frac{log_2(x_{ki}x_{kj})}{2}$$

3. Subtract the normalization curve fits  $M_k' = M_k - \hat{M}_k$  and obtain adjusted probe intensities:

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

 Take each of these adjustments (one for each pairwise comparison between arrays) and weight them equally across the set of arrays.

### Contrast method

Very similar to the cyclic loess method, because it's another way of normalizing based on M vs. A:

- 1. Data is converted to the log scale and the basis is transformed (this is just a fancy linear algebra step).
- 2. n-1 normalizing curves are fit to the transformed basis as in cyclic loess.
- Data is transformed again so that the normalizing curves lie on the horizontal, this time using a smooth function.
- 4. This normalized data is transformed back to the original basis and exponentiated.

This is slightly faster than cyclic loess but fitting the curves can still be slow.

### Contrast method

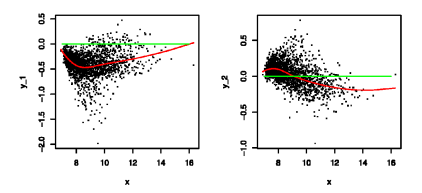


Figure 2: Scatter plots of the 2 contrasts against the mean for 3 arrays, A, B, and C, prior to normalizing. The red curve is the fitted normalizing curve, and the green line is the reference line. (Åstrand, 2003)

# The quantile method

- The goal of this method is to standardize the distribution of probe intensities across all arrays.
- ▶ The approach is an *n*-dimensional extension of the fact that, given a quantile-quantile plot where all of the points are on a straight diagonal line, you can be fairly sure that the two data vectors have the same distribution.
- We want to project our data onto the unit vector  $(\frac{1}{\sqrt{n}},...,\frac{1}{\sqrt{n}})$ .
- Let  $\mathbf{q}_k = (q_{k1}, ..., q_{kn})$  be the vector of  $k^{\text{th}}$  quantiles for k = 1, ..., p. Then:

$$proj_{\mathbf{d}}\mathbf{q}_{k} = (\frac{1}{n}\sum_{i=1}^{n}q_{kj},...,\frac{1}{n}\sum_{i=1}^{n}q_{kj})$$

## Quantile normalization algorithm

- 1. Given n arrays with p probe intensity measurements, make the  $p \times n$  matrix X, where each column has all the data from a single array.
- 2. Sort each column of X to produce  $X_{sort}$ . So, each row in  $X_{sort}$  is a quantile.
- 3. Take the mean of each row, and replace every value in the row with the mean to produce  $X'_{\rm sort}$
- 4. Put each column of  $X'_{sort}$  back in the original ordering from X to produce  $X_{normalized}$

This approach could theoretically be a problem for probes that have the same value across all arrays, but in practice this isn't an issue.

# Scaling method

Based on the approach suggested by Affymetrix, but this paper uses a probe-level version.

- 1. Choose a baseline array  $x_{\text{base}}$ : Usually this is the median array, but doesn't necessarily have to be.
- 2. For each other array, calculate the mean trimmed intensity  $\tilde{x_i}$  and find

$$\beta_{\it i} = \frac{\tilde{\it x}_{\rm base}}{\tilde{\it x}_{\it i}}$$

3. Normalized intensities are  $x_i' = \beta_i x_i$ 

### Non-linear method

- ▶ The scaling method is the same as fitting a straight line with intercept 0 between  $x_{\text{base}}$  and  $x_i$ .
- ► This can be extended to non-linear methods, usually a loess curve such that:

$$x_i' = \hat{f}_i(x_i)$$

where  $\hat{f}_i(\cdot)$  is the curve mapping from array i to baseline.

### Data

### Bolstad et al. tested these techniques on two datasets:

- 1. 30 arrays each from liver and central nervous system cell lines (6 groups at 5 dilution levels), plus 15 arrays with mixtures of cell lines in 75:25, 50:50, and 25:75 proportions.
- A dilution series of 27 arrays where 11 cDNA fragments have been spiked in at various concentrations. They also use two sets of triplicates (6 arrays total) from a Latin square experiment.

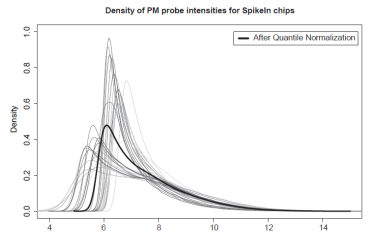


Figure 3: A plot of the densities for PM for each of the 27 spike-in datasets, with distribution after quantile normalization superimposed. (Bolstad et al., 2003)

log(PM)

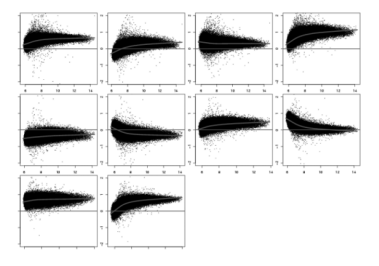


Figure 4: 10 pairwise M versus A plots using liver (at concentration 10) dilution series data for unadjusted data. (Bolstad et al., 2003)

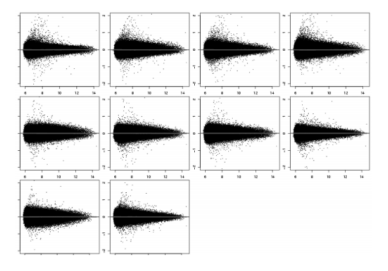


Figure 5: 10 pairwise M versus A plots using liver (at concentration 10) dilution series data after quantile normalization. (Bolstad et al., 2003)

- ➤ To compare methods at the probeset level, this paper uses the Robust Multichip Average (RMA).
- ▶ Plotting the log variance ratio against the log mean between two methods allows you to examine "differences in the between array variations and intensity dependent trends."
- When the loess line is below the x-axis, the first method has lower variance, and vice versa when the line is above the horizontal axis.

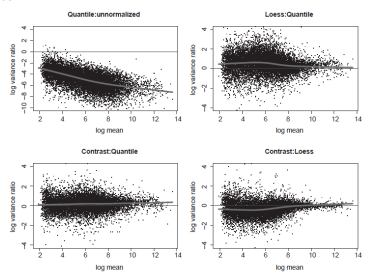


Figure 6:  $log_2$  variance ratio versus average  $log_2$  mean across 5 arrays for liver dilution data at concentration 10. (Bolstad et al., 2003)

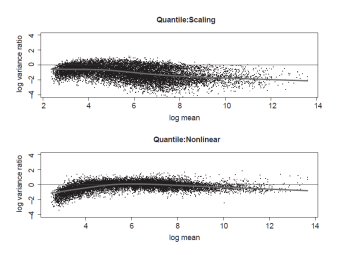


Figure 7:  $log_2$  variance ratio versus average  $log_2$  mean using the spike-in data. Comparing the baseline methods with the quantile method. (Bolstad et al., 2003)

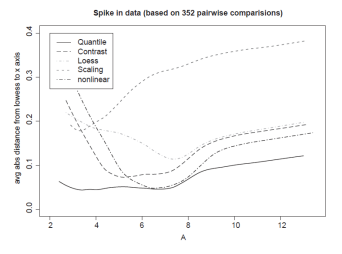


Figure 8: Comparing the ability of methods to reduce pairwise differences between arrays by using average absolute distance from loess smoother to x-axis in pairwise M versus A plots using spike-in dataset. Smaller distances are favorable. (Bolstad et al., 2003)

► The spike-in data can be used to assess bias for these techniques by fitting a regression model that includes the known concentration:

$$\log_2(\mathsf{Expression}) = \beta_0 + \beta_1 \log_2(\mathsf{concentration}) + \epsilon$$

- Ideally you would see  $\beta_1=1$  for the spike-in probesets and  $\beta_0=0$  for the non-spike-in sets.
- ► However, none of the median slopes for the non-spike-in probesets are exactly 0, which suggests that spike in concentration affects intensity.
- ➤ The authors recommend adjusting the slopes of the normalization procedures but subtracting the median slope of the non-spike-ins (e.g. 0.845 + 0.005 = 0.850).
- ► There is likely some error due to "pipette" effect, so concentrations may not be exactly correct.

Table 1. Regression slope estimates for spike-in probesets. A slope closer to one is better

Name	Quantile	Contrast	Loess	Non-linear	Scaling	None
AFFX-BioB-5_at	0.845	0.837	0.834	0.803	0.850	0.893
AFFX-DapX-M_at	0.778	0.771	0.770	0.746	0.783	0.826
AFFX-DapX-5_at	0.754	0.747	0.728	0.731	0.764	0.807
AFFX-CreX-5_at	0.903	0.897	0.889	0.875	0.912	0.955
AFFX-BioB-3_at	0.836	0.834	0.825	0.807	0.848	0.890
AFFX-BioB-M_at	0.789	0.782	0.781	0.762	0.797	0.838
AFFX-BioDn-3_at	0.547	0.543	0.550	0.514	0.553	0.595
AFFX-BioC-5_at	0.801	0.794	0.793	0.763	0.808	0.851
AFFX-BioC-3_at	0.796	0.790	0.785	0.769	0.805	0.847
AFFX-DapX-3_at	0.812	0.804	0.793	0.776	0.815	0.859
AFFX-CreX-3_at	-0.007	-0.006	0.002	-0.007	0.005	0.046
Non-spike-in (median)	-0.005	-0.005	-0.005	-0.007	-0.001	0.042

Figure 9: (Bolstad et al., 2003)

- Next they compared the means produced by quantile normalization and the non-linear method.
- ▶ Used two sets of triplicates (6 arrays) where the fold change between the spike-in concentrations is high.
- About half of the spike-in probesets are high in one triplicate and low in the other, and the opposite for the remaining probesets.
- For the non-linear method, they changed which array was used as baseline, including two "synthetic" baseline arrays made by taking probe-wise means and medians.

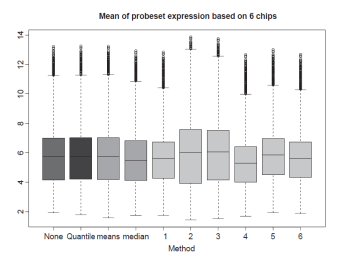


Figure 10: Distribution of average (over 6 chips) of a probeset expression measure using different baseline normalizations. (Bolstad et al., 2003)

Table 2. Comparing variance and bias with the non-linear normalization when using different baselines

Method	% with lower var reduced cf. U	% lower var reduced cf. Q	Abs Bias	# abs Bias cf U	# abs Bias cf Q
Probewise mean	83	40	9.2	5	5
Probewise median	96	58	7.9	6	6
Non-linear 1	96	53	7.5	7	5
Non-linear 2	93	31	11.8	2	4
Non-linear 3	94	37	10.5	4	4
Non-linear 4	95	47	7.4	6	5
Non-linear 5	96	55	7.4	7	5
Non-linear 6	96	55	7.5	7	5
Quantile (Q)	95	NA	8.5	6	NA
Unnormalized (U)	NA	NA	9.7	NA	NA

Figure 11: (Bolstad et al., 2003)

- ► For all approaches, about 95% of probesets have reduced variance.
- ➤ Compared to the quantile approach, over 50% of the probesets have reduced variance for 4 approaches (median, NL1, NL5, and NL6).

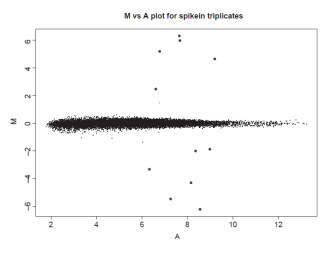


Figure 12: M versus A plot for spike-in triplicate data normalized using quantile normalization. Spike-ins are clearly identified. (Bolstad et al., 2003)

### Conclusions

- All three complete data methods (cyclic loess, contrast, and quantile) reduced variability across arrays more than scaling. The non-linear approached performed about as well as all three.
- ▶ The complete data methods performed comparably in terms of bias. The non-linear approach did not perform well in spike-in regressions. The slopes for scaling were closer to 1 but more variable than other methods.
- ► Choice of baseline can affect downstream analysis, so complete data methods are preferable.
- Quantile normalization is the fastest of the complete data methods, and therefore the best approach.

## Questions

- 1. What are some potential disadvantages of the quantile approach?
- 2. What about advantages of the Affymetrix-recommended approach?
- 3. What are some potential problems with this paper's methodology?
- 4. Would you draw the same conclusions as the authors based on these results?

### References

- Dudoit, S., Yang, Y. H., Callow, M. J., & Speed, T. P. (2002). Statistical Methods for Identifying Differentially Expressed Genes in Replicated cDNA Microarray Experiments. Statistica Sinica, 12(1), 111–139. JSTOR.
- Åstrand, M. (2003). Contrast Normalization of Oligonucleotide Arrays. Journal of Computational Biology, 10(1), 95–102. https://doi.org/10.1089/106652703763255697
- Bolstad, B. M., Irizarry, R. A., Åstrand, M., & Speed, T. P. (2003). A comparison of normalization methods for high density oligonucleotide array data based on variance and bias. Bioinformatics, 19(2), 185–193. https://doi.org/10.1093/bioinformatics/19.2.185