Normalization

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Biological vs. technical variability in gene expression

- What is ultimately of interest in the use of gene expression microarrays is the measurement of differences between experimental and reference states or between different groups of experimental units.
- Observed differences in microarray gene expression studies, however, are recognized as arising from two sources:
 - Biological variability changes in signal intensity driven by changes between biological states (healthy – disease)
 - Technical variability non-biological sources of variability

Sources of technical variability

Systematic

- Amount of extracted RNA, efficiencies of RNA extraction, reverse transcription, labeling, photodetection, GC content of probes
- · Similar non-bilological effect on many measurements
- Corrections can be estimated from data and accounted for by normalization

3/28

Sources of technical variability

Stochastic

- PCR yield, DNA quality, spotting efficiency, spot size, non-specific hybridization, stray signal
- Noise components & "Schmutz" (dirt)
- Too random to be explicitly accounted for need to use error modeling

Why normalization

Main idea

- Remove the systematic bias in the data as completely as possible while preserving the variation in the gene expression that occurs because of biologically relevant changes in transcription.
- The purpose of normalization is to adjust the gene expression values so that all genes on the array that are not differentially expressed have similar values across all arrays.

5/28

Goal of normalization

Assumption

- The average gene does not change in its expression level in the biological sample being tested.
- Most genes are not differentially expressed
- Up- and down-regulated genes roughly cancel out the expression effect.

Two categories of normalization methods

Baseline (reference) based methods

 Use a reference set of selected genes (housekeeping, invariant, spike-ins), or a baseline array

Complete (global, scaling) methods

· Combine information from all arrays in a given dataset

7/28

Reference set

- Housekeeping genes responsible for essential activities of cell maintenance & survival but not involved in cell function or proliferation. Such genes will be similarly expressed in all samples.
- Control genes serve as artificial housekeeping gene set that should have equal expression across arrays or channels

Reference set

- Invariant set genes that have the same rank across experiments.
 Empirically chosen
- All genes appropriate when the majority of the genes are believed to be not differentially expressed
- Problems defining reference sets may be biased. E.g., invariant set genes will be selected from the center of the distribution

9/28

Within- and between array normalization

Intra-slide normalization (within array)

- Applies to two-channel arrays
- Normalizes expression values to make intensities in two channels consistent within each array

Inter-slide normalization (between array)

- Normalizes expression values to achieve consistency between arrays
- · Generally done after within-array normalization

Normalization procedure

The normalized signal intensity ratio for clone k on array j will be

$$x_{jk} = \log \frac{R_{jk}}{G_{jk}} - c_{jk}$$

Where

- · R_{jk} the (background adjusted) Red signal
- · G_{jk} the (background adjusted) Green signal
- · c_{jk} the normalization factor

11/28

Calculating c_{jk}

- · Global normalization
- · Intensity dependent normalization
 - Lo(w)ess
 - Invariant set
 - Quantile

Global normalization

 c_{jk} is the same for all genes on array j.

Underlying assumptions

- Red & Green intensities have ~linear relationship through the origin;
- All cDNA species within a sample will incorporate an equivalent amount of dye per mole cDNA;
- There are no other variables that contribute to dye bias across slides.

13/28

Calculating c_{jk}

A constant c_j equal to the mean or median of the log ratios may be subtracted from all spots on array j. For example,

$$c_{jk} = c_j = median \left(log \frac{R_{jk}}{G_{jk}} \right)$$

for all clones/probes k in S.

Alternatively, fit a linear regression and use the estimated slope parameter as the constant.

Disadvantages of global normalization

- · Does not account for non-linearity of signal intensities.
- · Assumes cDNA from both dyes hybridized equally.
- More commonly, intensity dependent normalization methods are used.

15/28

Intensity-dependent normalization

Corrects intensities depending on the level of intensity, thereby changing the shape of the distribution of data

- · Bland Altman (MA) plots
- · Fitting a non-linear exponential curve
- · LOWESS/LOESS regression

Intensity dependent normalization

· Here the correction is still

$$x_{jk} = \left(log\frac{R_{jk}}{G_{jk}}\right) - c_{jk}$$

but now c_{jk} is the lowess fit, or $c_{jk} = f_j(A_{jk})$ where f is some smoothing function fitted to array j over all clones/probes k in S.

- Robust locally weighted regression of intensity log-ratios M_{jk} on the average log-intensity A_{jk} overall (global lowess) can be used for intensity dependent normalization.
- Other methods such as smoothing splines or exponential fits may also work well.

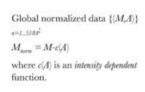
17/28

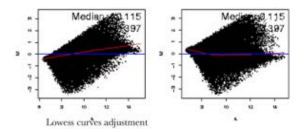
Intensity-dependent normalization: LOWESS

- LOcally WEighted Scatterplot Smoothing (Cleveland, 1979)
- First proposed for microarrays by Yang et al. (2002). Yang et al. (2002) used local window of 40%.
- Global LOWESS use implicit assumptions that, when stratified by mRNA abundance,
 - Only a minority of genes are expected to be differentially expressed or,
 - Any differential expression is as likely to be up-regulation as well as down-regulation

Intensity-dependent normalization: LOWESS

- · Loess normalization is based on MA plots.
- Skewing reflects experimental artifacts such as the contamination of one RNA source with genomic DNA or rRNA, the use of unequal amounts of fluorescent probes.



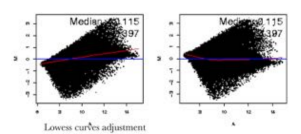


19/28

Intensity-dependent normalization: LOWESS

- Skewing can be corrected with local smoother: fitting a local regression curve to the data and subtracting the predicted value from the observed values
- Goal: minimize the standard deviation and place the mean log ratio at 0

Global normalized data $\{(M_rA)\}$ $n=1..5186^{\frac{1}{r}}$ $M_{norm}=M\text{-}c(A)$ where c(A) is an intensity dependent function.



Print-tip lowess

- · LOWESS fits to the data within print-tip groups
- Sub-array normalization

21/28

Affymetrix Method

- Scaling (Affymetrix method, <u>sadd_whitepaper</u>): First, choose a baseline GeneChip against which all other GeneChips are normalized.
- Calculate the 2% trimmed mean expression for the baseline GeneChip, represented by \widetilde{x}_{base} .
- Calculate the 2% trimmed mean expression for the j^{th} GeneChip, represented by \widetilde{x}_j .
- The scaling factor is taken to be $\beta_j=\widetilde{x}_{base}/\widetilde{x}_j$, so that the scaled values on GeneChip j are

$$x_{ik}^{scaled} = \beta_j * \widetilde{x}_{jk}$$

Rank invariant set

- Rather than using all genes for normalization, one may want to restrict the set of genes used for normalization by identifying those that are invariant.
- First, for each chip all genes are ranked; the invariant set is the set of genes with the same rank for each of the chips.
 - This is usually a very small number hence typically genes with approximately the same rank are used.
- Once the set of rank invariant genes is identified, intensity dependent normalization (fitting some smooth fit) is typically applied.

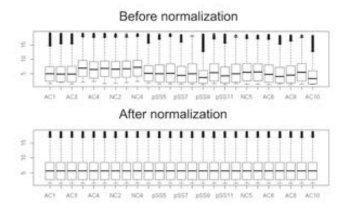
23/28

Quantile normalization

- Motivation from quantile-quantile plot
- Normal quantile-quantile plot consists of a plot of the ordered values in your data versus the corresponding quantiles of a standard normal distribution
- If the normal qqplot is fairly linear, your data are reasonably Gaussian; otherwise, they are not.

Between-array normalization methods

 Quantile normalization: Make distribution of data equal across all samples. Final distribution is the average of each quantile across chips (Bolstad et.al., Bioinformatics (2003))



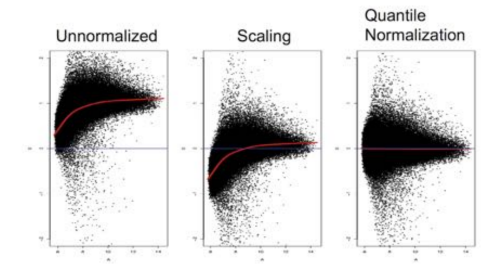
25/28

Quantile normalization

- 1. Given n arrays of length p, form matrix X of dimension $p \times n$ where each array is a column.
- 2. Sort each column of X to get X_{sort} . Remember to original order
- 3. Take the means across rows of X_{sort} and replace the values of X by those means. The resulting matrix is X'_{sort} .
- 4. Get $X_{normalized}$ by rearranging each column of X'_{sort} to have the same ordering as original X.

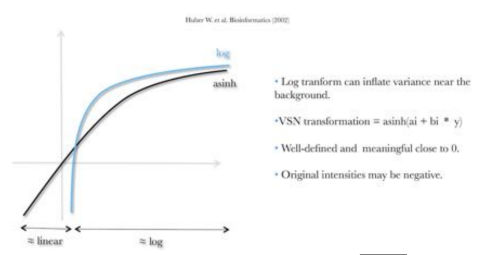
Quantile normalization changes expression over many slides i.e. changes the correlation structure of the data, may affect subsequent analysis.

Comparison of normalization techniques



27/28

Variance stabilizing normalization (VSN)



Inverse hyperbolic sine function $ashin\ x = ln(x + \sqrt{1 + x^2})$. Has the compressing effect on large values like regular $ln\ x$, but has much less of a compressing effect for small values. Defined on the entire real number line, no need to add an offset like for regular log-transformation.