Statistical Algorithms Description Document
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OVERVIEW

This document is a detailed reference guide for the *Statistical Algorithms* used in the analysis of GeneChip expression data. The guide focuses on how they work, what calculations and approaches they comprise, and how the tunable parameters are designed. Additional references are provided for additional information.

GENECHIP ARRAY DESIGN

It is important to understand how a GeneChip array is designed when considering the most appropriate approaches for its analysis. A GeneChip probe array consists of a number of probe cells where each probe cell contains a unique probe. Probes are tiled in probe pairs as a Perfect Match (PM) and a Mismatch (MM). The sequence for PM and MM are the same, except for a change to the Watson-Crick complement in the middle of the MM probe sequence. A probe set consists of a series of probe pairs and represents an expressed transcript.

DATA OUTPUTS

The *statistical algorithms* provide the following data outputs:

<table>
<thead>
<tr>
<th>Output</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Signal</td>
<td>A measure of the abundance of a transcript.</td>
</tr>
<tr>
<td>Stat Pairs</td>
<td>The number of probe pairs in the probe set.</td>
</tr>
<tr>
<td>Stat Pairs Used</td>
<td>The number of probe pairs in the probe set used in the Detection call.</td>
</tr>
<tr>
<td>Detection</td>
<td>Call indicating whether the transcript was Present (P) or Absent (A), or Marginal (M).</td>
</tr>
<tr>
<td>Detection p-value</td>
<td><em>p</em>-value indicating the significance of the Detection call.</td>
</tr>
<tr>
<td>Stat Common Pairs</td>
<td>The number of probe pairs in the probe sets from baseline and experimental arrays used in the Change call.</td>
</tr>
<tr>
<td>Change</td>
<td>Call indicating a change in transcript level between a baseline array and an experiment array [i.e. increase (I), decrease (D), marginal increase (MI), marginal decrease (MD), no change (NC)].</td>
</tr>
<tr>
<td>Change p-value</td>
<td><em>p</em>-value indicating the significance of the Change call.</td>
</tr>
<tr>
<td>Signal Log Ratio</td>
<td>The change in expression level for a transcript between a baseline and an experiment array. This change is expressed as the log₂ ratio. A log₂ ratio of 1 is the same as a Fold Change of 2.</td>
</tr>
<tr>
<td>Signal Log Ratio Low</td>
<td>The lower limit of the log ratio within a 95% confidence interval.</td>
</tr>
<tr>
<td>Signal Log Ratio High</td>
<td>The upper limit of the log ratio within a 95% confidence interval.</td>
</tr>
</tbody>
</table>
DATA PREPARATION
In this section we will discuss steps that occur prior to application of the new statistical algorithms.

A note about .CEL Files
The Statistical Algorithms begin with information contained in the .CEL file generated by Microarray Suite software. The .CEL files contain a captured image of the scanned GeneChip® array and calculations of the raw intensities for probe sets. The method for calculating individual cell intensities, thus generating the .CEL file, is not affected by the Statistical Algorithms. Therefore, it will not be discussed here.

Masking
Masked probe pairs are excluded from all algorithms. For more information about “probe masking” see the Affymetrix® GeneChip® Expression Analysis Technical Manual (2001), Section 4.1.17.

Background Subtraction
A calculated background establishes a “floor” to be subtracted from each cell value.

Zone Values
For purposes of calculating background values, the array is split up into $K$ rectangular zones $Z_k$ ($k = 1, \ldots, K$, default $K = 16$).
Control cells and masked cells are not used in the calculation.
The cells are ranked and the lowest 2% is chosen as the background $b$ for that zone ($b_{Z_k}$).
The standard deviation of the lowest 2% cell intensities is calculated as an estimate of the background variability $n$ for each zone ($n_{Z_k}$).

Smoothing Adjustment
To provide a smooth transition between zones, we compute distances from each cell on the chip to the various zone centers. A weighted sum is then calculated based on the reciprocal of a constant plus the square of the distances to all the zone centers. In more detail, if the distance $d$ between the chip coordinate $(x, y)$ and the center of the $k^{th}$ zone is $d_k$, we can calculate a weighting factor, which is related to the square of $d$ (the relationship between $w$ and $d$ is illustrated in the graph next page). A small factor, smooth, is added to $d^2$ to ensure that the value will never be zero.

A GeneChip® array is divided into a number of equally spaced zones and an average background is assigned to the center of the zone, indicated by green circles. For each cell, the distance is calculated to the center of every zone. A weighting factor is then calculated as the reciprocal of the sum of a constant and the square of the distance. The colors of the arrows indicate the relative weights.
\[ w_k(x, y) = \frac{1}{d_k^2(x, y) + \text{smooth}} \]

(default \(\text{smooth} = 100\))

For every cell, we can use the sum of all the weights to all the zone centers to calculate the background \(b\) value to be used for cell \(x, y\):

\[ b(x, y) = \frac{1}{\sum_{k=1}^{K} w_k(x, y) b_{zk}} \]

**Noise Correction**

Now we want to compute an adjusted value that shifts intensities down by the local background. In order to do so, we must first ensure that the values would not become negative. Negative intensity values are problematic later in the calculations when log values are calculated.

For noise correction, a local noise value \(n\) based on the standard deviation of the lowest 2% of the background in that zone \((n_{Z_k})\) is calculated and weighted for background values (just substitute \(n(x, y)\) for \(b(x, y)\) and \(n_{Z_k}\) for \(b_{Z_k}\) in the formula above).

Then a threshold and a floor are set at some fraction of the local noise value, so that no value is adjusted below that threshold. That is, for a cell intensity \(I'(x, y)\) at chip coordinates \((x, y)\), we compute an adjusted intensity.

\[
A(x, y) = \max(I'(x, y) - b(x, y), \text{NoiseFrac}\times n(x, y))
\]

where \(I'(x, y) = \max(I'(x, y), 0.5)\)

NoiseFrac is the selected fraction of the global background variation.

(default \(\text{NoiseFrac} = 0.5\))
EXPRESSION VALUE CALCULATION (SIGNAL)

The Signal value is calculated from the combined, background-adjusted, PM and MM values of the probe set. It represents the amount of transcript in solution.

Signal is calculated as follows:
1. Cell intensities are preprocessed for global background.
2. An ideal mismatch value is calculated and subtracted to adjust the PM intensity.
3. The adjusted PM intensities are log-transformed to stabilize the variance.
4. The biweight estimator (see Appendix I) is used to provide a robust mean of the resulting values. Signal is output as the antilog of the resulting value.
5. Finally, Signal is scaled using a trimmed mean.

Method

Background and Contrast Correction

Before we can proceed, we need to do the background subtraction as described in the Data Preparation section.

Ideal Mismatch (IM)

The reason for including a MM probe is to provide a value that comprises most of the background cross-hybridization and stray signal affecting the PM probe. It also contains a portion of the true target signal. If the MM value is less than the PM value, it is a physically possible estimate for background, and can be directly used.

If the MM value is larger than the PM value, it is a physically impossible estimate for the amount of stray signal in the PM intensity. Instead, an idealized value can be estimated based on our knowledge of the whole probe set or on the behavior of probes in general. Specifically, we base this estimate either on the average ratio between PM and MM, or (if that measure is itself too small) a value slightly smaller than PM.

To calculate a specific background ratio representative for the probe set, we use the one-step biweight algorithm ($T_{bi}$), which is described in Appendix I. We find a typical log ratio of PM to MM that is simply an estimate of the difference of log intensities for a selected probe set. The biweight specific background ($SB$) for probe pair $j$ in probe set $i$ is:

$$SB_i = T_{bi} \left( \log_2 (PM_{i,j}) - \log_2 (MM_{i,j}) : j = 1, \ldots, n_i \right)$$

Throughout this text we use log base 2 exclusively.
If $SB_i$ is large, then the values from the probe set are generally reliable, and we can use $SB_i$ to construct the ideal mismatch $IM$ for a probe pair if needed. If $SB_i$ is small ($SB_i \leq \text{contrast } \tau$), we smoothly degrade to use more of the PM value as the ideal mismatch. The three cases of determining ideal mismatch $IM$ for probe pair $j$ in probe set $i$ are described in the following formula:

$$IM_{i,j} = \begin{cases} 
MM_{i,j}, & \text{if } MM_{i,j} < PM_{i,j} \\
\frac{PM_{i,j}}{2^{(SB_i)}}, & \text{if } MM_{i,j} \geq PM_{i,j} \text{ and } SB_i > \text{contrast } \tau \\
\frac{PM_{i,j}}{2^\left(\frac{\text{contrast } \tau}{\text{scale}_{i}}\right)}, & \text{if } MM_{i,j} \geq PM_{i,j} \text{ and } SB_i \leq \text{contrast } \tau
\end{cases}$$

default $\text{contrast } \tau=0.03$

default $\text{scale } \tau = 10$

The first case where the mismatch value provides a probe-specific estimate of stray signal is the best situation. In the second case, the estimate is not probe-specific, but at least provides information specific to the probe set. The third case involves the least informative estimate, based only weakly on probe-set specific data.

The blue bars represent PM and the red bars MM probes of a hypothetical probe set. In the top panel, most of the MM values are smaller than PM, so we use the MM directly. The yellow bars indicate the estimated value we would use if MM>PM.

For probe pair 4, the MM is larger than the PM, so it is not a useful value for estimating the stray signal component of PM. An imperfect, but useful resolution is to estimate a MM value that is typical for the probe set. For the overall probe set, the mean difference $SB_i$ of the logs of PM and MM is large, so we can use it directly to estimate a value for MM 4. The yellow bar indicates the estimate.

In the second panel, $SB_i$ is small, so we cannot base an accurate estimate for MM 4 on it. The best we can do is to calculate a value (indicated by the yellow bar) slightly less than PM.
Probe Value and Signal Log Value
Given the ideal mismatch value, the formula for the probe value (PV) is fairly simple. To guarantee numerical stability, we use the formula:
\[
V_{i,j} = \max\left( PM_{i,j} - IM_{i,j}, \delta \right) \text{ default } \delta = 2^{-20}
\]

Now we calculate the probe value PV for every probe pair \( j \) in probeset \( i \). \( n \) is the number of probe pairs in the probeset.
\[
PV_{i,j} = \log_2 (V_{i,j}),\ j = 1, \ldots, n_i
\]

We then compute the absolute expression value for probe set \( i \) as the one-step biweight estimate (see Appendix I) of the \( i \ n \) adjusted probe values:
\[
\text{SignalLogValue}_i = T_{bi}(PV_{i,1}, \ldots, PV_{i,n})
\]

Scaled Probe Value
Note: the scaling \( (sf) \) and normalization factors \( (nf) \) computed in this section are reported by the software.

If the algorithm settings indicate scaling all probes sets or selected probe sets to a target we calculate a scaling factor \( (sf) \)
\[
sf = \frac{Sc}{\text{TrimMean}\left(\text{SignalLogValue}_i, 0.02, 0.98\right)}
\]
where \( Sc \) is the target signal (default \( Sc = 500 \)) and the SignalLogValues in the SignalLogValue, set are the probe sets indicated in the algorithm settings. The \text{TrimMean} function here takes the average value of all observations after removing the values in the lowest 2\% of observations and removing those values in the upper 2\% of observations. If the algorithm settings indicate user defined scaling, then \( sf = \) user defined value.

The reported value of probe set \( i \) is:
\[
\text{ReportedValue}(i) = nf \times sf \times 2^{(\text{SignalLogValue}_i)}
\]
where \( nf = 1 \) for absolute analysis and is computed as follows for a comparison analysis.

If the algorithm settings indicate user defined normalization, then \( nf = \) user specified normalization.

Otherwise, the algorithm settings either indicate normalizing all or selected probe sets:
\[
nf = \frac{\text{TrimMean}\left(\text{SPV}_b, 0.02, 0.98\right)}{\text{TrimMean}\left(\text{SPV}_e, 0.02, 0.98\right)}
\]
where \( \text{SPV}_b \) is the baseline signal, and \( \text{SPV}_e \) is the experiment signal (scaled-only) and \( i \) defines the probe sets selected by the user.

This is reported as Signal.
Since comparison analyses are done on matched probe pairs, the individual probe pair values are also modified by this scaling factor. The scaled probe value $SPV_{i,j}$ is

$$SPV_{i,j} = PV_{i,j} + \log_2 (nf \times sf)$$

These values are used in computing the log-ratio in comparison analysis.

**Performance on corrupted data**

The algorithms were developed on a very clean data set, where every precaution was taken to ensure that the data was of high quality. It is important to validate the performance on poor quality data. Good performance on poor quality data will be an indication of how well the algorithms will perform under real-world situations, where the data is less than perfect. To generate data with a controlled amount of noise, several .CEL files were used and an increasing amount of data was replaced with random numbers. The random numbers were chosen uniformly within the typical intensity range (0 to 46,000). The correlation coefficient between the true concentration and the output from the newly generated corrupted data set was then plotted as a function of the proportion of uncorrupted data. When the proportion of uncorrupted data is 1 (the original data set), we see a correlation coefficient near 1.0 (even the original data set has some noise). As the proportion of good data decreases, we expect the predictive power of the data with respect to concentration to decrease. This loss of predictive power is reflected by a lower correlation coefficient.

![Performance of methods against bad data (human)](corrupt data chosen from uniform[0…46000])

Performance against bad data (human). This graph illustrates the robustness against bad data obtained by using this analysis strategy. The data in several .CEL files from the human array HG-U95A are degraded by substituting random numbers between 0 and 46,000. The intact data set is indicated by 1 on the x-axis. As the amount of intact data decreases, the average values start deviating from the original. The data from the Signal Calculation Algorithm implemented in MAS 4.0 also starts deviating, while the biweight algorithm remains accurate even after 20% of the data were corrupted.
For averages (blue line) the correlation coefficient drops rapidly indicating that the results quickly become less accurate. This is not surprising, because an average is not robust against outliers. The AvgDiff algorithm used in MAS 4.0 (pink line) is more robust because it discards some outliers. However, this strategy loses power when the data is increasingly corrupt, because it has only a small ability to identify outliers when much of the data is corrupt. The Signal algorithm implemented in MAS 5.0 is very robust against corrupted data and the results remain well-correlated even when as much as 20% of the data is corrupt. Naturally, additional noise never improves the quality of the data, and so does degrade the results, but robustness provides a safety net against corrupted data completely destroying the utility of an array.
RELATIVE EXPRESSION VALUES (LOG RATIOS)

By doing a direct cell to cell comparison between probe sets on two arrays, systematic probe effects will be canceled out. Probe effects refer to the inherent differences in the hybridization efficiency of different probes, which is a source of variation in sampling signals from different sequences present at the same concentration, even when physically linked in the same nucleic acid polymer. Calculating the ratio of signal for the same probe on two different arrays effectively cancels the intrinsic affinity factor for that sequence.

**Adjusted cell intensities (Baseline and Experiment) ➠ Log Ratio, Log Ratio Low and Log Ratio High**

**Method**
The Signal Log Ratio calculation is an extension of the Signal calculation.
- Scale the baseline and experiment.
- Correct for probe pair bias.
- Calculate the signal log ratio.

**Nomenclature**

Comparative experiments are always run on two arrays, one assigned baseline \( b \) and the other experiment \( e \). Here we show the same probeset \( i \) from two arrays.

**Signal Log Ratio**

Once we have a scaled probe value each probe pair (SPV is calculated in the previous section), we can calculate the Signal log ratio using the biweight algorithm (see Appendix I). The probe log ratio \( PLR \) is calculated for probe pair \( j \) in probeset \( i \) on both the baseline \( b \) and experiment \( e \) arrays:

\[
PLR_{i,j} = e^{SPV_{i,j} - bSPV_{i,j}}
\]
If we have the probe log ratios PLR we can use the biweight algorithm to calculate the SignalLogRatio (see Appendix I for a description of the biweight algorithm).

$$\text{SignalLogRatio}_i = T_{bi}(PLR_{i,1}, \ldots, PLR_{i,n_i})$$

From the biweight calculation, we can also determine the 95\textsuperscript{th} confidence interval and report the results as Log Ratio Low and Log Ratio High. Thus, the relative expression for a target in two samples being compared is estimated by calculating the average of the log (ratios) for each corresponding probe pair in the probe sets.

**Fold Change**

Previous versions of the Affymetrix\textsuperscript{®} Microarray Suite software communicated the relative signal level between the same probe set on two different arrays as a fold-change ratio that was signed depending on the direction of the change in an ordered pair. It is straightforward to convert from the LogRatio metric to the older Fold-Change value.

Fold Change from log ratios is calculated as follows:

Corrected formula

$$
FoldChange_i = \begin{cases} 
2^{\text{SignalLogRatio}_i} & \text{SignalLogRatio}_i \geq 0 \\
(-1) \times 2^{-\text{SignalLogRatio}_i} & \text{SignalLogRatio}_i < 0 
\end{cases}
$$
SINGLE ARRAY ANALYSIS (DETECTION CALLS)

A detection call answers the question: “Is the transcript of a particular gene Present or Absent?” In this context, Absent means **that the expression level is below the threshold of detection.** That is, the expression level is not provably different from zero. In the case of an uncertainty, we can get a Marginal call. *It is important to note that some probe-sets are more variable than others,* and the minimal expression level provably different from zero may range from a small value to very large value (for a noisy probe-set). The advantage to asking the question in this way without actual expression values is that the results are easy to filter and easy to interpret. For example, we can imagine that we may only want to look at genes whose transcripts are detectable in a particular experiment.

**Raw cell intensities ➤ Absent, Present, or Marginal call plus p-values**

**Method**

There are four steps to the method:

1. Remove saturated probe pairs and ignore probe pairs wherein PM ~ MM + tau
2. Calculate the discrimination scores. (This tells us how different the PM and MM cells are.)
3. Use Wilcoxon’s rank test to calculate a significance or p-value. (This tells us how confident we can be about a certain result.)
4. Compare the p-value with our preset significance levels to make the call.

**Saturation**

If a mismatch cell is saturated \( MM \geq 46000 \), the corresponding probe pair is not used in further computations. We also discard pairs where PM and MM are within \( \tau \) of each other.

If all probe pairs in a unit are saturated, we report the gene as detected and set the \( p \)-value to 0.

The statistical significance, or \( p \)-value, of a result is the probability that the observation in a sample occurred merely by chance under the null hypothesis. The null hypothesis is that the target is absent (zero effect on the probes). For example, a \( p \)-value of 0.005 means that less than 5 out of 1000 probe sets for absent transcripts will be called present based on the distribution of intensity within the corresponding probe sets that is equally or less likely occur by chance. In detection, the smaller the \( p \)-value, the more significant the results suggesting that the gene may be present.

![Graph showing discrimination score and MM intensity/probe pair]

In this hypothetical probe set the PM intensity is 80 and the MM intensity for each probe pair increases from 0 to 100. The discrimination score offers a smooth function that decreases as the MM intensity increases. In other words, as the intensity of the MM increases our ability to discriminate between the PM and MM decreases. Note that the value becomes negative when \( MM > PM \).

The broken line indicates the threshold .
Discrimination Score

The discrimination score \([R]\) is a relative measure of the difference between the PM and MM intensities. The discrimination score for the \(i\)th probe pair is:

\[
R_i = \frac{PM_i - MM_i}{PM_i + MM_i}
\]

We use \(0.02\) (default), a small threshold between 0 and 1 as a small significant difference from zero. If the median \((R_i) > 0\), we can reject the hypothesis that PM and MM are equally hybridizing to the sample. We can make a detection call based on the strength of this rejection (the \(p\)-value).

Increasing the threshold can reduce the number of false detected calls, but may also reduce the number of true detected calls.

Computing \(p\)-values:

The one-sided Wilcoxon’s Signed Rank Test (Appendix II) is used in the Call algorithms and is used to calculate the \(p\)-values for the null hypothesis:

\(H_0: median(R_i - \tau) = 0\) versus the alternative hypothesis:

\(H_1: median(R_i - \tau) > 0\)

Note: detection calls are calculated on the raw intensity values, so prevents false detected calls where PM is only slightly larger than MM.

Note: There is a relationship between the discrimination scores and the \(\log_2\) ratio used in the Specific Background (SB) calculation. It is known as “Fisher’s z-transformation.”

\[
\begin{align*}
\text{log}_2 \frac{1+r}{1-r} &= \log_2 \frac{PM-MM}{PM+MM} \\
&= \log_2 \frac{1+\frac{PM-MM}{PM+MM}}{1-\frac{PM-MM}{PM+MM}} \\
&= \log_2 \frac{PM+MM+PM-MM}{PM-MM} \\
&= \log_2 \left( \frac{PM+MM}{2*PM} \right) - \log_2 \left( \frac{PM-MM}{2*MM} \right)
\end{align*}
\]

HINT: See Appendix for complete description of the one-sided Wilcoxon’s Signed Rank Test calculation.

Determining the default. The discrimination scores of 8960 probe pairs spiked at known concentrations on the human U95Av2 array and yeast S98 array were examined. The default value was selected to fall between the discrimination ratio of spikes at zero and 0.25pM concentration.
Making the call

We set two significance levels \( \alpha_1 \) and \( \alpha_2 \) such that
\[
0 < \alpha_1 < \alpha_2 < 0.5
\]

\( \alpha_{\text{default}} = 0.04 \) (16-20 probe pairs)
\( \alpha_{\text{default}} = 0.06 \) (16-20 probe pairs)

Reducing the significance level \( \alpha_1 \) can reduce the number of false Detected calls and reduce the number of true Detected calls.

Increasing the significance level \( \alpha_2 \) can reduce the number of false undetected calls and reduce the number of true undetected calls.

\( (*) \) Although in an ideal world, the distribution of probes exactly matches the statistical assumptions, in practical cases the distribution is not precise. Our thresholds therefore do not result in the exact false-positive rate specified by the alpha parameter when the target is absent. This rate will vary with extraneous factors, including the number of probe-pairs used.
COMPARATIVE ANALYSIS (COMPARISON CALLS)
A comparative call answers the question: “Does the expression level of a transcript on one chip (experiment) change (significantly) with respect to the other chip (baseline)?” The possible distinct answers are Increase, Marginally Increased, No Change detected, Marginally Decreased, and Decrease. As with detection calls, No Change means the difference is below the threshold of detection. That is, the difference is not provably different from zero. It is important to note that some probe-sets are more variable than others, and the minimal expression difference provably different from zero may range from a small value to very large value (for a noisy probe-set, or for low concentrations). Note that saturated probe pairs are excluded from the computation of comparative calls. If all probe pairs of a probe set are saturated, we report that no comparative call can be made.

We attempt to find changes in expression level by examining changes in the intensities of both PM and MM probes between experiments. The differences in PM and MM in both experiments and differences between PM and background in both experiments are examined using a non-parametric Wilcoxon rank test to look for significant differences.

Differences between PM and MM and Differences between PM and Background (Experimental and Baseline) ➤ Change call and $p$-value

Saturation
If one of the four cells (PM and MM in Baseline and PM and MM in Experiment) is saturated ($PM_i$ or $MM_i$ = 46000), the corresponding probe pair is not used in further computations. The number of discarded cells can be determined from the Stat Common Pairs parameter.

Quantities Used in Comparative Calls
For a probe set of $n$ probe pairs, we form two $n$-dimensional vectors for comparative calls.

$$q = (q_1, \ldots, q_n) \text{ and } z = (z_1, \ldots, z_n)$$

The component $q_i$ is the difference between Perfect Match intensity $PM_i$ and Mismatch intensity $MM_i$ for the $i^{th}$ probe pair:

$$q_i = PM_i - MM_i, \ (i = 1, \ldots, n),$$

and the component $z_i$ is the difference between the Perfect Match intensity $PM_i$ and background level $b_i$:

$$z_i = PM_i - b_i, \ (i = 1, \ldots, n)$$
$b$ is the background and is calculated the same as for Data Preparation ($b(x,y)$).

Using both $q$ and $z$ can produce better empirical call results than using only one of them.

**Balancing Factors**

Note: The factors computed in this section are not reported by the software.

The distributions of $q$ and $z$ over all probe pairs in an experiment are slightly and subtly different from each other, and are different between two experiments. We therefore provide a balancing factor for each type of data to correct some of this difference. (The distributions of $q$ and $z$ are also not identical to the Signal distribution, and therefore the scaling factor used for signal is not used here).

Vectors $q$ and $z$ have two different balancing factors:

$e[i]$ is a modified average of $q_j$ for the probe pairs of transcript $i$, and is calculated as the average of all $q$ values within three standard deviations from the average $q$ for transcript $i$. We calculate a global balancing factor using the trimmed mean of $e[i]$ over all transcripts $i$:

$$sf = \frac{Sc}{\text{trimmedMean}(e[i],0.02,0.98)}$$

If $\text{trimmedMean}(e[i],0.02,0.98) \leq 0$, the scaling factor can be calculated as:

$$sf = \frac{Sc}{\text{trimmedMean}(\max(e[i],0),0.02,0.98)}$$

These values are calculated for both the experiment ($E$) and the baseline ($B$) arrays. Now the primary balancing factor $nf$ can be calculated:

$$nf = \frac{sf_E}{sf_B}$$

A second, primary balancing factor $nf_2$ is calculated for $z$. The calculations are exactly the same as for $q$, except only $z$ values are used. These values are calculated for both the experiment ($E$) and the baseline ($B$) arrays to give:

$$nf_2 = \frac{sf_2E}{sf_2B}$$

The two balancing factors are combined to match the distributions of $q$ and $z$ over the whole signal range.

However, any calculated balancing factor is only an approximation to the true differences between the distributions. To allow for small differences between the distributions not covered by the balancing factor, we will use a range of balancing factors. We straddle the
true balancing function by using three different balancing factors $f[k]$ for $q$, as well as three different balancing factors $f_2[k]$ for $z$ ($k = 0, 1, 2$). They are defined as:

$$
\begin{align*}
    f[0] &= nf \times d \\
    f[1] &= nf \\
    f[2] &= \frac{nf}{d} \\
    f_2[0] &=nf_2 \times d \\
    f_2[1] &= nf_2 \\
    f_2[2] &= \frac{nf_2}{d}
\end{align*}
$$

where $d \leq 1$, (default = 1.1).

**Special Case**

If the algorithm settings indicate a user defined balancing factor and the factor is not equal to 1 then, $nf = nf_2 = user\ defined\ normalization\ factor \times sfE / sfB$

where $sfE$ is the experiment $sf$ and $sfB$ is the baseline $sf$ as described in the Scaled Probe Value section.

---

**Wilcoxon Signed Rank Test (Appendix I)**

For every unit, we can form the $(2n)$-dimensional vector $qB$ in the baseline and vector $qE$ in the experiment. For $k = 0, 1, 2$, three vectors were formed:

$$
\begin{align*}
    v[k][i] &= f[k] \times qE[i] - qB[i] \\
    v[k][i \times n] &= C \times (f_2[k] \times zE[i] - zB[i]) \\
    (i &= 1, \ldots, n; \ k = 0, 1, 2) \\
    \text{default } C &= 0.2
\end{align*}
$$

and three, one-sided $p$-values $p[k]$ from the signed rank tests of the null hypothesis

$$
H_0 : \text{median}(v[k]) = 0
$$

versus the alternative hypothesis

$$
H_1 : \text{median}(v[k]) > 0.
$$

We use significance levels $1$ and $2$ to make calls. $1$ is the small significance level for comparative calls of a unit (an interpolation of $1H$ and $1L$), and is 0.0025 for default values of $1H$ and $1L$. $2$, the large significance level for comparative calls of a unit (an interpolation of $2H$ and $2L$), and is 0.003 for default values of $2H$ and $2L$. $1$ and $2$ should satisfy the relationship $0 < 1L < 2L < 0.5$ and $0 < 1H < 2H < 0.5$.

The $1$ value is a linear interpolation of $1L$ and $1H$. Similarly $2$ is derived from $2L$ and $2H$. The ability to adjust the stringency of calls associated with high and low signal ranges independently makes it possible to compensate for effects that influence calls based on low and high signals. However, this feature is not used by default, because the defaults are set as $1L = 1H$ and $2L = 2H$. 
The cut off $p$-values may appear very small if you are used to the usual significance value of 0.05. This is due to two reasons:

Due to the multiple measurement problem, we require smaller $p$-values to ensure that we do not get too many false calls for the whole collection of transcripts. This is similar to a Bonferroni correction, but we determine the value empirically.

The $p$-value produced by this calculation is also an over estimate of significance, because we are using two values, PM-MM and PM-background, that are not truly independent measures. Further, the null distribution does not exactly describe the empirical situation, and so critical thresholds may change with the number of probes.

However, the purpose of the $p$-values produced here is to rank results in order of significance, so the absolute $p$-value is not important.

We use the “critical” $p$-value as our output $p$-value.

The critical $p$-value, $p$, is the most conservative in making increase and decrease calls. It is defined by the following formula:

\[
p = \begin{cases} 
\max(p_0, p_1, p_2) & \text{if } p_0 < 0.5, p_1 < 0.5 \text{ and } p_2 < 0.5 \\
\min(p_0, p_1, p_2) & \text{if } p_0 > 0.5, p_1 > 0.5 \text{ and } p_2 > 0.5 \\
0.5 & \text{otherwise}
\end{cases}
\]
The margins can also be represented as follows (16-20 probe pairs):

<table>
<thead>
<tr>
<th>Increase</th>
<th>Adjusting Parameters</th>
</tr>
</thead>
</table>
| \begin{align*} \text{increase} & : \{p[0] < \gamma_1, \quad p[1] < \gamma_1, \quad p[2] < \gamma_1\} \\
| \text{marginally increase but not increase} & : \{p[0] < \gamma_2, \quad p[1] < \gamma_2, \quad p[2] < \gamma_2\} \\
| \text{decrease} & : \{p[0] > 1 - \gamma_1, \quad p[1] > 1 - \gamma_1, \quad p[2] > 1 - \gamma_1\} \\
| \text{marginally decrease but not decrease} & : \{p[0] > 1 - \gamma_2, \quad p[1] > 1 - \gamma_2, \quad p[2] > 1 - \gamma_2\} \\

If none of the above conditions are satisfied, we make a No-Change call.

**Adjusting Parameters**

- Decreasing \((\_L \text{ and } \_H)\) can reduce the number of false Increase and Decrease calls, but can also reduce the number of true Increase and Decrease calls.
- Increasing \((\_L \text{ and } \_H)\) can reduce the number of false No-Change-detected calls, but can also reduce the number of true No-Change-detected calls.
- Increasing the perturbation parameter \(d\) can increase the number of true No-Change-detected calls, but can also increase the number of false No-change-detected calls.

It is recommended to use two or more identical sample replicates to adjust parameters.

All comparative call thresholds should ideally be empirically determined from the false-positive rate. Set an allowed false change call rate, e.g., 0.01. Sort the \(p\)-values and find the significant level that give this rate of increasing or decreasing calls. Do not set the allowed false change call rate too small, because it will be very difficult to make increasing or decreasing calls for other experiments where these calls are reasonable.
REFERENCES


Wilcoxon, F. Individual Comparisons by Ranking Methods *Biometrics* 1, 80-83 (1945).

APPENDIX I

One-Step Tukey’s Biweight Algorithm

Purpose
There are several stages in the algorithms in which we want to calculate an average. The biweight algorithm is a method to determine a robust average unaffected by outliers.

1. First, the median is determined to define the center of the data.
2. Then the distance of each data point from the median is determined. This distance is then used to determine how much each value should contribute to the average. For example, outliers that are far away from the median should contribute less to the average.

The full biweight algorithm iterates through a process of calculating the estimate and then reweighting the observations until there is no further change. We found that the first step of the biweight iteration provides the most useful increase in quality.

Method
The one-step biweight algorithm begins by calculating the median $M$ for a data set with $n$ values. In the Signal measurement, this data set consists of the log (PM-IM) probe values of a probe set with $n$ probe pairs.

Next, we calculate the absolute distance for each data point from the median. We then calculate $S$, the median of the absolute distances from $M$. The Median Absolute Deviation, $MAD$, is an initial measure of spread.

For each data point $i$, a uniform measure of distance from the center is given by:

$$u_i = \frac{x_i - M}{cS + \varepsilon}, i = 1, \ldots, n$$

$c$ is a tuning constant (default $c = 5$).

$\varepsilon$ is a very small value used to avoid zero values in the division (default $\varepsilon = 0.0001$).
Weights are then calculated by the bisquare function:

\[
   w(u) = \begin{cases} 
   (1-u^2)^2, & |u| \leq 1 \\
   0, & |u| > 1 
   \end{cases}
\]

For each point, the weight \( w \) is reduced by a function of its distance from the median so outliers are effectively discounted by a smooth function. When values are very far from the median, their weights are reduced to zero.

The corrected values can now be calculated by using the one-step \( w \)-estimate (a weighted mean):

\[
   T_{bi} = \frac{\sum w(u) x_i}{\sum w(u)}
\]

“The performance of one-step-from-the-median \( w \)-estimators is essentially as good as that of fully iterated M-estimates” (Hoaglin, Mosteller, Tukey). We follow their suggestion and save computation by using this \( w \)-estimate.

The \( t \) distribution can then be used to determine a confidence interval length.

**Confidence Intervals**

An additional benefit of the biweight algorithm is that we can calculate confidence limits using standard statistics.

The first step is to calculate the measure of uncertainty for the biweight \( s_{bi} \):

\[
   s_{bi} = \sqrt{n} \sqrt{\frac{\sum_{i=1}^{n} (x_i - T_{bi})^2 (1-u_i^2)^4}{\sum_{i=1}^{n} (1-u_i^2)(1-5u_i^2)}}
\]

An approximate 95% confidence interval is then computed as

\[
   T_{bi}(x_1, x_2, \ldots, x_n) \pm t_{df}^{(0.975)} \frac{s_{bi}}{\sqrt{n}}
\]

where \( t_{df}^{(0.975)} \) is the 97.5\(^{th}\) percentile for the \( t \) distribution with the number of degrees of freedom set equal to \( \max(0.7 \cdot (n - 1), 1) \).
APPENDIX II

One-sided Wilcoxon’s Signed Rank Test

Wilcoxon’s signed rank test possesses many good properties of nonparametric statistical methods.
1. It does not assume a normal data distribution.
2. It is robust and insensitive to outliers.
3. It can be used on raw data without much handling.

The signed rank test applies to two paired data sets. For example PMi and MMi probes in a probe set or (PMi-MMi)Baseline and (PMi-MMi)Experiment from paired data sets. To demonstrate signed rank test procedure, we consider two paired data sets: baseline b and experiment e with n probe pairs, b = (b1, ..., bn) and e = (e1, ..., en) and calculate di = ei - bi for every probe pair.

Uninformative (Tied) Probe Pairs

We first calculate the absolute differences |di| for all pairs of data. We exclude the probe pairs whose di = 0 from further computation. If all differences are zero, we output 0.5 as the one-sided p-value.

Rank Sum

The steps for calculating the rank sum for non-zero differences are as follows:

<table>
<thead>
<tr>
<th>Start values</th>
<th>Convert to absolute values</th>
<th>Sort in increasing order</th>
<th>Assign ranks</th>
<th>Place ranks back in their original order</th>
<th>Their signed ranks</th>
</tr>
</thead>
<tbody>
<tr>
<td>di = 2</td>
<td></td>
<td>a4 = 0.5</td>
<td>r4 = 1.5</td>
<td></td>
<td>S4 = 4.5</td>
</tr>
<tr>
<td>di = 1</td>
<td></td>
<td>a3 = 0.5</td>
<td>r3 = 1.5</td>
<td></td>
<td>S3 = 3</td>
</tr>
<tr>
<td>di = -2</td>
<td></td>
<td>a2 = 1</td>
<td>r2 = 3</td>
<td></td>
<td>S2 = -4.5</td>
</tr>
<tr>
<td>di = 0.5</td>
<td></td>
<td>a1 = 2</td>
<td>r1 = 4.5</td>
<td>S1 = 4.5</td>
<td>S1 = -4.5</td>
</tr>
<tr>
<td>di = 0.5</td>
<td></td>
<td>a0 = 0.5</td>
<td>r0 = 4.5</td>
<td></td>
<td>S0 = 1.5</td>
</tr>
</tbody>
</table>

Then we form the sum of positive signed ranks. In our example, S = s1 + s2 + s4 + s5 = 10.5.

NOTE: we use the one-sided test here. For the one-sided test, if the null hypothesis is true, the p-value should be uniformly distributed between 0 and 1. In other words, if there is no difference between the experiment and the baseline the p-value should be “near” 0.5.

When the alternative hypothesis is true (i.e., there is a positive change), the p-value should be close to 0.

When median (g, h) < 0 is true, the p-value should be close to 1. This property makes the one-sided test useful for both absolute and comparative calls.

NOTE: where values are identical, such as R4 and R5 the values are ranked 1 and 2 and we assign the average, which is 1.5, to both of them.
Confidence Values
The changes in probe set values may vary greatly, from very small to very large. For example, our confidence in the change will be low if the changes are small or close to background. Large, significant changes will have high confidence. The goal is to be able to set a confidence level and give the final call based on whether or not the confidence level exceeds a certain threshold. This will allow us to only accept calls in which we have confidence, at the risk of missing small changes that may be real, but have a low confidence threshold. Conversely, we may be interested in finding as many changes as possible at the risk of including changes that may be accidental.

Fortunately, an advantage of the one-sided Wilcoxon’s Signed Rank Test is that there are well-known methods for calculating $p$-values. The formulas are fairly complex, but not critical to understanding how the basic method works. We use two different methods: one for large probe sets and one for small probe sets.

Small Probe Sets
When the number of probe pairs $n$ is small ($n < 12$), we can simply enumerate all the possible outcomes and compute the $p$-value directly. In this case, we apply signs to ranks $r_i$ ($i = 1, 2, \ldots, n$) in every possible way, calculate the sum of positive ranks and denote this sum by $S_j$ ($j = 1, \ldots, 2^n$).

$$ p(S) = 2^{-n} \sum_{j=1}^{2^n} u(S_j > S) + 0.5u(S_j = S) $$

where $u()$ is the characteristic function; when the argument is a logical expression, it is one if the argument is true, and it is zero if the argument is false; when the argument is a numeric expression, it is one if the argument is positive, and it is zero otherwise. Since these assignments of ranks are equally probable, we simply need to count the number of instances in which they are as large as our observed value.

In our example, all possible signed ranks and the sum of positive ranks $S_j$ are listed (see table). Since the order of these ranks does not matter, we use the ascending order of their absolute values in the table and denote them by $s'_j$. 
Random Signed Ranks for p-value Evaluation

<table>
<thead>
<tr>
<th>J</th>
<th>s'₁</th>
<th>s'₂</th>
<th>s'₃</th>
<th>s'₄</th>
<th>s'₅</th>
<th>S'_{J}</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1.5</td>
<td>-1.5</td>
<td>-3</td>
<td>-4.5</td>
<td>-4.5</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>1.5</td>
<td>-1.5</td>
<td>3</td>
<td>-4.5</td>
<td>-4.5</td>
<td>1</td>
</tr>
<tr>
<td>3</td>
<td>-1.5</td>
<td>1.5</td>
<td>-3</td>
<td>-4.5</td>
<td>-4.5</td>
<td>1</td>
</tr>
<tr>
<td>4</td>
<td>-1.5</td>
<td>1.5</td>
<td>3</td>
<td>-4.5</td>
<td>-4.5</td>
<td>1</td>
</tr>
<tr>
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<td>1.5</td>
<td>-3</td>
<td>4.5</td>
<td>-4.5</td>
<td>4.5</td>
</tr>
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<td>3</td>
<td>-4.5</td>
<td>4.5</td>
<td>4.5</td>
</tr>
<tr>
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<td>-1.5</td>
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<td>-4.5</td>
<td>4.5</td>
<td>3</td>
</tr>
<tr>
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<td>-1.5</td>
<td>3</td>
<td>-4.5</td>
<td>4.5</td>
<td>4.5</td>
</tr>
<tr>
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<td>1.5</td>
<td>-1.5</td>
<td>-3</td>
<td>4.5</td>
<td>-4.5</td>
<td>6</td>
</tr>
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<td>4.5</td>
<td>6</td>
</tr>
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<td>1.5</td>
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<td>-4.5</td>
<td>4.5</td>
<td>6</td>
</tr>
<tr>
<td>12</td>
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<td>1.5</td>
<td>-3</td>
<td>-4.5</td>
<td>4.5</td>
<td>6</td>
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<td>4.5</td>
<td>-4.5</td>
<td>9</td>
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<td>9</td>
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<td>4.5</td>
<td>9</td>
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<td>4.5</td>
<td>4.5</td>
<td>9</td>
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</tr>
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<td>4.5</td>
<td>4.5</td>
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</tr>
<tr>
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<td>1.5</td>
<td>-3</td>
<td>4.5</td>
<td>4.5</td>
<td>10.5</td>
</tr>
<tr>
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<td>4.5</td>
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</tr>
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<td>-3</td>
<td>4.5</td>
<td>4.5</td>
<td>12</td>
</tr>
<tr>
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<td>3</td>
<td>4.5</td>
<td>4.5</td>
<td>12</td>
</tr>
<tr>
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<td>1.5</td>
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<td>4.5</td>
<td>4.5</td>
<td>13.5</td>
</tr>
<tr>
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<td>1.5</td>
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<td>4.5</td>
<td>4.5</td>
<td>13.5</td>
</tr>
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<td>4.5</td>
<td>13.5</td>
</tr>
<tr>
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<td>1.5</td>
<td>3</td>
<td>4.5</td>
<td>4.5</td>
<td>15</td>
</tr>
</tbody>
</table>

The table shows the 32 possible results for the five probe ranks shown (2⁵ possible ways of assigning signs to each rank).

All signed ranks above 10.5 are given a weight of 1 (there are five in Table 1) and items with signed ranks equal 10.5 are given a rank of .5 (there are four in Table 1).

In our example  \( p(10.5) = \frac{(1*5) + (0.5*4)}{32} = 0.21875 \)

Large Probe Sets
When the number of probe pairs \( n \) is large (in our implementation, \( n \geq 12 \)), we use the asymptotic approximation. The statistic \( S' \) is considered to have a standard normal distribution with mean 0 and variance 1, where:

\[
S' = \frac{S - \frac{n(n+1)}{4}}{\sqrt{\frac{n(n+1)}{24} - \nu t}}
\]

where \( \nu t \) is a term modifying the variance for ties. The formula for \( \nu t \) is:

\[
\nu t = \sum_{k=1}^{t} \frac{b_k (b_k^2 - 1)}{48}, \text{ for } k = 1 : t
\]

where \( t \) is the number of tie groups; \( b_k \) is the number of ties in the tie group \( k \). The one-sided p-value is

\[
p(S) = 1 - f(S')
\]

where \( f(S') \) is the standard normal cumulative distribution function.

NOTE: Under certain circumstances “Fisher's permutation test” may have more power than Wilcoxon rank. We elected to use the same test across all numbers of probes to be consistent, even if it has lower power than the best possible test.
APPENDIX III

Noise (Q) Calculation

The calculation of the Q value is provided here for completeness. It is not used anywhere in the statistical algorithm. However, since it is based on pixels it provides a useful quality measure of how well the grid was placed on the array to calculate the .CEL file. All other calculations are derived from the .CEL file.

Q, the noise for a given probe array hybridization, is calculated by taking the average (over all the cells used in background computation) of the following value in each cell: standard deviation of the pixel intensity (st devi) divided by the square root of the pixel number (\( \sqrt{\text{pixel}_i} \)):

\[
Q = \frac{1}{N} \left( \sum_{i} \frac{\text{stdev}_i}{\sqrt{\text{pixel}_i}} \right) \times SF \times NF
\]

where \( N \) is the total number of background cells for an array, \( \text{stdev}_i \) is the standard deviation of the intensities of the pixels making up feature \( i \), \( \text{pixel}_i \) is the number of pixels in feature \( i \), \( SF \) is the scaling factor, and \( NF \) is the normalization factor for the analysis.

NOTE: The noise (Q) is calculated from pixel values in the DAT file.