

# Selection and validation of normalization methods for c-DNA microarrays using within-array replications

Jianqing Fan\* and Yue Niu

Department of Operations Research and Financial Engineering  
Princeton University, Princeton, NJ 08544

## ABSTRACT

**Motivation:** Normalization of microarray data is essential for multiple-array analyses. Several normalization protocols have been proposed based on different biological or statistical assumptions. A fundamental problem arises whether they have effectively normalized arrays. In addition, for a given array, the question arises how to choose a method to most effectively normalize the microarray data.

**Results:** We propose several techniques to compare the effectiveness of different normalization methods. We approach the problem by constructing statistics to test whether there are any systematic biases in the expression profiles among duplicated spots within an array. The test statistics involve estimating the genewise variances. This is accomplished by using several novel methods, including empirical Bayes methods for moderating the genewise variances and the smoothing methods for aggregating variance information. P-values are estimated based on a normal or chi-square approximation. With estimated P-values, we can choose a most appropriate method to normalize a specific array and assess the extent to which the systematic biases due to the variations of experimental conditions have been removed. The effectiveness and validity of the proposed methods are convincingly illustrated by a carefully designed simulation study. The method is further illustrated by an application to human placenta cDNAs comprising a large number of clones with replications, a customized microarray experiment carrying just a few hundred genes on the study of the molecular roles of Interferons on tumor, and the Agilent microarrays carrying tens of thousands of total RNA samples in the MAQC project on the study of reproducibility, sensitivity and specificity of the data.

**Availability:** Code to implement the method in the statistical package R is available from the authors.

**Contact:** jqfan@princeton.edu

## 1 INTRODUCTION

Microarray techniques have been widely used in many areas of biological research. They have substantial impact on tumor diagnosis, and classification and understanding of the molecular mechanisms of biochemical processes, tumorigenesis and tumor developments. Proper statistical analysis is vital for revealing meaningful biological results. For an overview of statistical analysis of DNA microarrays, we refer to Fan and Ren 2006 and references therein.

The quality of microarray data is very important for downstream statistical analysis (Marshall, 2004; Eisenstein, 2006; Shi *et al.*,

2006, Patterson *et al.*, 2006). Experimental variations, such as RNA quality, probe labeling, hybridization condition, washing, signal and background detection in the scanning process, slide and block effects, pose significant challenges in the analysis of microarray data. The first step in microarray analysis is to remove the systematic biases due to the variations in experimental conditions so as to make multiple array analyses meaningful. These efforts are collectively referred to as the normalization of microarray data in the literature.

A number of useful normalization protocols have been proposed based on different assumptions. These include the global normalization (Kroll and Wolf, 2002), rank invariant normalization (Tseng *et al.*, 2001), LOWESS normalization (Dudoit *et al.*, 2002), Semi-Linear In-slide Model (SLIM, Fan *et al.*, 2004, 2005), Two-way Semi-Linear Models (TW-SLM, Huang *et al.*, 2005), and robust TW-SLM (Wang *et al.*, 2005; Ma *et al.*, 2006). Recently, two seminal normalization methods (CADS and eCADS) based on dye swap and statistical models are proposed by Dabney and Storey (2007a, b). All of the aforementioned methods are based on some statistical and biological assumptions. For example, the global normalization is adequate only when there is no print-tip block effect and no intensity effect; the LOWESS method assumes implicitly that at each given intensive level the average expression level of up- and down-regulated genes are about the same in each print-tip block; SLIM, TW-SLM and robust TW-SLM all require that the statistical models are correct. The questions then arise naturally for a given array, which methods are most effective to normalize the data and whether the data have been properly normalized. These questions are very fundamental to the statistical analysis of microarray and have not yet been addressed.

Our study is motivated by the aforementioned fundamental concerns. Our approach is to use the duplicated spots within an array. They contain the most valuable information about possible systematic biases in the microarray experiments. The idea is that if microarray data have been properly normalized, there should be no systematic biases among duplicated spots. Therefore, when the sum of the square differences of duplicated spots, standardized by the estimated genewise variances, are aggregated among many different genes having duplications, the test statistic follows approximately a  $\chi^2$ -distribution with a large degree of freedom, or more formally a normal distribution. This provides a simple and useful diagnostic test statistic to check if an array has been properly normalized by a particular method. Regarding the test statistic as a measure of the discrepancy of replicated spots after normalization, we select a normalization method that has the smallest value of the test statistics. In addition, the associated P-value of the test statistic

\*to whom correspondence should be addressed

enables us to judge the degree to which the normalization has been properly carried out.

In implementing the validation tests, it involves inevitably the estimation of genewise standard deviations (SD) and variances. A precise estimate of standard deviations and variances will improve the statistical power of the validation tests. It has also important applications in selecting significant genes (Tusher *et al.*, 2001; Dudoit *et al.*, 2003; Reiner *et al.*, 2003; Storey and Tibshirani, 2003; Cui *et al.*, 2005; Smyth *et al.*, 2005). The precision of estimating genewise SDs and variances depends on the number of replications that are available. Several innovative strategies will be introduced to enhance the precision of the estimate.

Duplicated spots play very important roles in the analysis of microarray data. They are not only powerful for normalization (Fan *et al.*, 2004), but also useful for genewise variance estimation (Smyth *et al.*, 2005) in selecting statistically differently expressed genes. Furthermore, they are fundamental to our proposed validation tests. The availability of such duplicated genes can be accomplished by the designs of c-DNA microarrays. For example, in the microarrays analyzed by Fan *et al.* 2004, 111 out of 19968 clones of genes were printed twice randomly on the 32 print-tip blocks. This enables them to untangle the block effects and intensity effects from these 111 duplicated spots. With the increased popularity of customized microarrays, which enables researchers to focus only on hundreds of genes of their primary interest with more reliable measurements, within-array replications can easily be obtained. The gene selection biases in customized arrays require more sophisticated normalization techniques. The validation tests are essential for controlling the quality of downstream statistical data analysis of customized arrays.

## 2 METHODS OF NORMALIZATION

There are several useful normalization methods, which are based on different biological or statistical assumptions. We briefly review several of them that will be used in our numerical studies.

Suppose that we have  $J$  replications of a c-DNA microarray experiment. For each given array, there are  $N$  different genes. Among them,  $G$  genes are replicated  $I$  times. Let  $I_g$  denote the number of replications for gene  $g$ , with  $I_g = 1$  indicating no duplication. For customized arrays, usually  $G = N$  (all genes have within-array replications) and  $I_g = 2$  or 3 or 4. However, this can also be designed differently. For the microarrays analyzed in Fan *et al.* 2004,  $G = 111$  and  $N = 19968 - G$  so that  $I_g = 2$  only for 111 clones that are duplicated and randomly placed on 32 print-tip blocks.

Let  $R_{gij}$  and  $G_{gij}$  be respectively the intensities of red (Cy3) and green (Cy5) channels for the  $i^{th}$  duplication of the  $g^{th}$  gene in the  $j^{th}$  array, and  $b_{gi}$  be the print-tip block where the  $i^{th}$  duplication of the  $g^{th}$  gene resides. Then, we can compute the log-ratios and log-indensities as

$$Y_{gij} = \log_2(G_{gij}/R_{gij}), \quad \text{and} \quad X_{gij} = 0.5 \log_2(G_{gij} * R_{gij}).$$

The global normalization is to compute the median  $\hat{m}_j$  of log-ratios  $\{Y_{gij}\}$  of the  $j^{th}$  array and to normalize the data for the  $j^{th}$  array as

$$\hat{Y}_{gij} = Y_{gij} - \hat{m}_j. \quad (1)$$

This basically assumes that the up-regulated and down-regulated genes are about the same, which does not usually hold for

customized arrays due to gene selection biases. In addition, the global normalization assumes no block or intensity effects.

To address the above two concerns, Dudoit *et al.*, 2002 apply the global normalization technique more locally to each block and each intensity level, resulting in the LOWESS normalization. For the data  $\{(X_{gij}, Y_{gij}) : b_{gi} = b\}$  in a given print-tip block  $b$  of the  $j^{th}$  array, they applied the LOWESS smoother to estimate the conditional mean function  $\hat{m}_{bj}(x)$  of the log-ratios  $Y$  given the intensity level  $X = x$ , and computed the normalized log-ratios in the  $b^{th}$  block in  $j^{th}$  array as follows:

$$\hat{Y}_{gij} = Y_{gij} - \hat{m}_{bj}(X_{gij}), \quad \text{with } b_{gi} = b. \quad (2)$$

This significantly relaxes the restrictions of the global normalization, but still assumes that up-regulated and down-regulated genes are about the same at each given intensity level. Again, this might not be appropriate when the cells or tissues are treated by cytokines. It might not be valid for customized arrays due to gene selection biases.

To address these issues, Fan *et al.*, 2004 introduced the SLIM technique based on the model assumption:

$$Y_{gij} = \alpha_g + \beta_{j,b_{gi}} + m_j(X_{gij}) + \varepsilon_{gij}, \quad (3)$$

in which  $\alpha_g$  is the treatment effect on gene  $g$ ,  $\beta_j$  and  $m_j$  represent respectively the array-specific block and intensity effect, and  $\varepsilon$  is the stochastic noise. For each given array  $j$ , Fan *et al.*, 2004 estimated the model parameters using the data from duplicated spots and computed the normalized data

$$\hat{Y}_{gij} = Y_{gij} - \hat{\beta}_{j,b_{gi}} - \hat{m}_j(X_{gij}). \quad (4)$$

In Fan *et al.*, 2005, the efficiency of parameters in (4) is improved by aggregating the data from all arrays. Namely, the parameters in (3) are jointly estimated using all arrays.

TW-SLM (Huang *et al.*, 2005) and robust TW-SLM (Wang *et al.*, 2005; Ma *et al.*, 2006) are in the same spirit as SLIM. It does not require duplications of spots but relies heavily on the assumption that the treatment effect  $\alpha_g$  is independent of the array. Statistically, it is a specific model of (3) with  $\beta = 0$ . With estimated parameters, the normalized data are computed as in (4) with  $\hat{\beta}_{j,b_{gi}} = 0$ . TW-SLM applies the technique for each given block so that the block effects in each array have been properly taken care of.

The effectiveness of SLIM, TW-SLM and robust TW-SLM depends critically on the statistical model assumption. With so many normalization methods, the questions arise naturally which method is the most appropriate for a specific array and whether the expression profiles have been properly normalized.

The CADS and eCADS (Dabney and Storey, 2007a, b) are the normalization methods based on dye-swap and statistical models. They aim at preserving time differential expression relationships among the treatment and control arrays after normalization.

## 3 VALIDATION TESTS

Within array replications not only provide useful information about the possible systematic biases: block effect and intensity effect, but also play an important role in validation tests of the necessity and effectiveness of the normalization methods. When the log-ratios

have been properly normalized, the systematic biases should be negligible and hence the following model holds approximately:

$$\hat{Y}_{gij} = \alpha_g + \varepsilon_{gij}, \quad (5)$$

$$g = 1, \dots, N, \quad i = 1, \dots, I_g, \quad j = 1, \dots, J,$$

where  $I_g$  is the number of duplication for gene  $g$ . See also (4).

Duplicated spots provide valuable information about the validity of model (5) for each given array. We will use only  $G$  genes with  $I$  replications for each given array to develop the validation tests, and drop the subscript  $j$  to facilitate the notation. This leads to the simplified notation:

$$\hat{Y}_{gi} = \alpha_g + \varepsilon_{gi}, \quad g = 1, \dots, G, \quad i = 1, \dots, I. \quad (6)$$

Hence, the difference  $\hat{Y}_{gi} - \bar{Y}_g$  should have mean zero, where  $\bar{Y}_g = \sum_{i=1}^I \hat{Y}_{gi}/I$ . We will assume that the first four moments of the noise behave like those of a normal distribution:

$$E\varepsilon_{gi} = 0, \quad E\varepsilon_{gi}^2 = \sigma_g^2, \quad E\varepsilon_{gi}^3 = 0, \quad E\varepsilon_{gi}^4 = 3\sigma_g^4.$$

### 3.1 Genewise standardization

Two natural test statistics for testing model (6) are based on the genewise standardized  $L_2$ - and  $L_1$ -norm, aggregated over  $G$  genes having duplications. Specifically, we let

$$T_1 = \sum_{g=1}^G \left\{ \sum_{i=1}^I (\hat{Y}_{gi} - \bar{Y}_g)^2 / \sigma_g^2 \right\}. \quad (7)$$

Under the normality assumption  $\varepsilon_{gi} \sim N(0, \sigma_g^2)$ , if the data have been properly normalized, the test statistic  $T_1$  follows the  $\chi^2$ -distribution with degree of freedom  $(I-1)G$ . In particular, when  $I = 2$ ,

$$T_1 = \sum_{g=1}^G (\hat{Y}_{g1} - \hat{Y}_{g2})^2 / (2\sigma_g^2) \sim \chi_G^2.$$

Note that the test statistic above is reasonably robust to the normality assumption. In the validation test, the number of genes with duplications  $G$  is reasonably large and by the Central Limit Theorem,  $T_1$  follows approximately a normal distribution. In this case,  $\chi_{(I-1)G}^2$  is also approximately a normal distribution. Hence, the null distribution  $\chi_{(I-1)G}^2$  is approximately valid.

Instead of using the  $L_2$ -norm, one can use a more robustified norm  $L_1$ -norm. This leads naturally to consider the test based on the genewise standardized  $L_1$ -norm:

$$T_2 = \sum_{g=1}^G \left\{ \sum_{i=1}^I |\hat{Y}_{gi} - \bar{Y}_g| / \sigma_g \right\}. \quad (8)$$

In particular, when  $I = 2$ , the test statistic reduces to

$$T_2 = \sum_{g=1}^G |\hat{Y}_{g1} - \hat{Y}_{g2}| / \sigma_g.$$

By the Central Limit Theorem,

$$T_2 \stackrel{\mathcal{L}}{\sim} N(\xi_I G, \eta_I^2 G), \quad (9)$$

where  $\xi_I = \sqrt{2I(I-1)/\pi}$  and  $\eta_I^2 = \text{var}(\sum_{i=1}^I |\hat{Y}_{gi} - \bar{Y}_g| / \sigma_g)$ . Specifically,

$$\eta_I^2 = \begin{cases} 2(1 - 2/\pi) = 0.7268, & \text{when } I = 2 \\ (4\sqrt{3} - 12)/\pi + 8/3 = 1.0523, & \text{when } I = 3. \end{cases}$$

### 3.2 Aggregated standardization

Accurate estimates of the genewise standard deviation  $\sigma_g$  are challenging. The process itself may depend on the selection of a method of normalization. The test statistics  $T_1$  and  $T_2$  may not be well approximated by the  $\chi^2$ -distribution or the normal distribution, when the genewise variances are not estimated accurately. In addition, the standardized square differences in  $T_1$  and  $T_2$  are sensitive to the estimation error in  $\sigma_g$ . On the other hand, the aggregated variance  $G^{-1} \sum_{g=1}^G \sigma_g^2$  or aggregated standard deviation  $G^{-1} \sum_{g=1}^G \sigma_g$  can be more accurately estimated. These considerations lead us to the unweighted differences among the expression profiles of replicated spots.

The aggregated differences among replicated spots when standardized, are given by

$$T_3 = \frac{\sum_{g=1}^G \sum_{i=1}^I (\hat{Y}_{gi} - \bar{Y}_g)^2 - (I-1) \sum_{g=1}^G \sigma_g^2}{\sqrt{2(I-1) \sum_{g=1}^G \sigma_g^4}}. \quad (10)$$

The test statistic  $T_3$  is obtained by aggregation first followed by standardization. On the other hand, the test statistic  $T_1$  is obtained by standardization first and followed by aggregation. The null distribution of  $T_3$  follows approximately  $N(0, 1)$  when  $G$  is large.

Following the same spirit, a more robustified counterpart of  $T_2$  is

$$T_4 = \left\{ \sum_{g=1}^G \sum_{i=1}^I |\hat{Y}_{gi} - \bar{Y}_g| - \xi_I \sum_{g=1}^G \sigma_g \right\} / \left\{ \eta_I \left( \sum_{g=1}^G \sigma_g^2 \right)^{1/2} \right\}. \quad (11)$$

where  $\xi_I$  and  $\eta_I$  are two constants defined in (9). The null distribution of  $T_4$  follows approximately  $N(0, 1)$ , when the data are properly normalized.

Note that the test statistic  $T_4$  involves the estimation of aggregated standard deviation  $\sum_{g=1}^G \sigma_g$ . If this is estimated with bias, then the null distribution will be shifted. The genewise variance is usually estimated by the sample variance or aggregated sample variance  $S_g^2$ . Suppose that  $S_g^2$  has the distribution  $K S_g^2 / \sigma_g^2 \sim \chi_K^2$  with a given degree of freedom  $K$ , then  $S_g^2$  is an unbiased estimator of  $\sigma_g^2$ , but  $S_g$  is not an unbiased estimator of  $\sigma_g$ . An unbiased estimator of  $\sigma_g$  is given by (Curland and Tripathi, 1971)

$$\{(K/2)^{1/2} \Gamma(K/2) / \Gamma((K+1)/2)\} S_g. \quad (12)$$

In our numerical studies, this is implemented in  $T_2$  and  $T_4$ . Our experiences show that the correction factor in (12) is necessary in order to obtain a more accurate approximation of the null distribution.

### 3.3 Choosing a method of normalization

The test statistics  $T_1, \dots, T_4$  can be regarded as measures of effectiveness of normalization. The smaller, the less discrepancy among repeated measurements, and the more effectiveness of a normalization method. For a given array, among several normalization methods, we would choose the one that has the smallest test statistic. The associated P-value gives us an idea on the extent to which the expression profiles have been normalized. The power of the validation test depends on the number of data points  $G$  in the training set. Excessively large  $G$  will result in overpower of the tests to reject even a tiny systematic bias.

As to be discussed in Section 4, the implementation of the validation tests might require the choice of an appropriate normalization method first in order to estimate the genewise variances. The aggregated standardization tests  $T_3$  and  $T_4$  can be used for this purpose, since the normalization constants can be ignored.

### 3.4 Training and testing sets

In many situations, there are many genes that have duplications. This is particularly the case for the customized arrays. In these situations, we can randomly select between 50 and 100 different genes as the testing set and use the remaining genes as the training set. The training set is used to estimate the parameters in the normalization, while the testing set is applied to the validation tests.

In other situations, there are limited genes that have duplicated spots. In this case, the multi-fold cross-validation ideas can be employed to choose the training and testing sets.

## 4 ESTIMATION OF GENEWISE VARIANCE

Accurate estimation of genewise variance  $\sigma_g^2$  is important for assessing the effectiveness of normalization. It is also critically important for selecting the genes that are statistically differently expressed among treatments and controls. See, for example, Tusher *et al.*, 2001, Storey and Tibshirani, 2003, Fan *et al.*, 2004, Cui *et al.*, 2005, and Fan and Ren 2006. In particular, Cui *et al.*(2005) demonstrates that genewise variance estimation has direct impact on the sensitivity and specificity of selecting differently expressed genes.

### 4.1 Use of within array replications

A natural estimate of the genewise variance is the sample variance of the duplicated expressions. These log-ratios are computed after the data are normalized. If there are several normalization methods available, one can use  $T_3$  and  $T_4$  without standardization to help select a method of normalization.

Suppose that we have  $J$  replicated arrays. If we assume that  $\text{var}(\hat{Y}_{gij}) = \sigma_g^2$ , which are the same across  $J$  arrays, then we would pool the variability information from other arrays. This leads to a pooled estimator of  $\sigma_g^2$  by

$$s_{W,g}^2 = \frac{1}{J(I-1)} \sum_{j=1}^J \sum_{i=1}^I (\hat{Y}_{gij} - \bar{Y}_{gj})^2. \quad (13)$$

The above estimate ignores the correlation among duplicated genes. If within-array replications have a common correlation  $\rho_g = \rho$  and observations across arrays are independent, Smyth *et al.*, 2005 introduced the residual maximum likelihood (REML) estimator of  $\sigma_g^2$  as follows:

$$s_g^2 = \frac{1}{IJ-1} \left\{ \frac{(J-1)s_{B,g}^2}{1+(I-1)\hat{\rho}} + \frac{J(I-1)s_{W,g}^2}{1-\hat{\rho}} \right\}, \quad (14)$$

where  $s_{B,g}^2 = I \sum_{j=1}^J (\bar{Y}_{gj} - \bar{Y}_g)^2 / (J-1)$  with  $\bar{Y}_g = \sum_{j=1}^J \bar{Y}_{gj} / J$  is the between-arrays variance and  $\hat{\rho}$  is an estimate of  $\rho$ . They also proposed the REML estimation of  $\hat{\rho}$  by

$$\hat{\rho} = \frac{\sum_{g=1}^G s_{B,g}^2 - \sum_{g=1}^G s_{W,g}^2}{\sum_{g=1}^G s_{B,g}^2 + (I-1) \sum_{g=1}^G s_{W,g}^2}.$$

They argued further that the degree of freedom of  $s_g^2$  is  $(IJ-1)$ .

### 4.2 Smoothing estimator

The degree of freedom for the within-array estimate  $s_{W,g}^2$  of the variance is still limited. As the variability of c-DNA microarray measurements is related to the intensity level (Tseng *et al.*, 2001, Fan *et al.*, 2004), one can pool the information of variability from expression profiles with similar intensity levels (Fan *et al.*, 2004). This results in a nonparametric estimate of the intensity-specific variance function  $\sigma^2(\cdot)$  from the nonparametric regression model:

$$\hat{Y}_{gi} = \alpha_g + \sigma(X_{gi})\eta_{gi}, \quad i = 1, \dots, I_g, \quad g = 1, \dots, N. \quad (15)$$

See also (5) with the array index  $j$  suppressed. The procedure of Fan *et al.*, 2004 is to first smooth  $\{\hat{Y}_{gi}\}$  on  $\{X_{gi}\}$  by using a local linear estimate, which is really a smoothing estimator of the scatter plot  $\{(X_{gi}, \hat{Y}_{gi})\}$ , to obtain an estimate of  $\alpha_g$  by regarding it as a smooth function of the log-intensity  $X_{gi}$ , and then smooth the squared residuals  $\{(\hat{Y}_{gi} - \hat{\alpha}_g)^2\}$  on  $\{X_{gi}\}$  to obtain an estimate of the intensity-specific variance function  $\sigma^2(\cdot)$ . The fundamental assumption in Fan *et al.*, 2004 is that  $\alpha_g$  is basically a smooth function of  $X_{gi}$ . When this assumption is not valid, the estimator will be biased.

The bias issue in Fan *et al.*, 2004 can be significantly reduced when the within-array replications are available as in (15). Consider those genes with  $I$  replications. Replacing  $\alpha_g$  by its unbiased estimate  $\bar{Y}_g = \sum_{i=1}^I \hat{Y}_{gi} / I$ , we have the variance of the residual

$$E(\hat{Y}_{gi} - \bar{Y}_g)^2 = (I-1)^2 \sigma^2(X_{gi}) / I^2 + \sum_{j \neq i} \sigma^2(X_{gj}) / I^2.$$

When the variability of log-intensities is not large, we can approximate  $X_{gj}$  by its average  $\bar{X}_g$ . Owing to the smoothness assumption, we have

$$\text{var}(\hat{Y}_{gi} - \bar{Y}_g) \approx (I-1)\sigma^2(\bar{X}_g) / I. \quad (16)$$

Letting  $r_g^2 = \sum_{i=1}^I (\hat{Y}_{gi} - \bar{Y}_g)^2 / (I-1)$  be the sample standard deviation, we have

$$Er_g^2 \approx \sigma^2(\bar{X}_g).$$

Therefore, the intensity-specific variance  $\sigma^2(\cdot)$  can be estimated by smoothing the pairs  $\{(\bar{X}_g, r_g^2) : g = 1, \dots, G\}$ , resulting in an estimated function  $\hat{\sigma}^2(\cdot)$ . Hence, our estimate of  $\sigma_g^2$  is

$$\hat{s}_g^2 = \hat{\sigma}^2(\bar{X}_g).$$

The approach is particularly appealing to the customized arrays, in which the variability of intensities is not large and the replicated spots are available. See Section 5.3.

### 4.3 Empirical Bayes estimator

With within-array replications, we have two estimators: REML  $s_g^2$  and the intensity-specific estimator  $\hat{s}_g^2$ . One naturally uses the intensity-specific estimator  $\hat{s}_g^2$  to augment the REML estimator  $s_g^2$ . One simple way to do this is the following empirical Bayes shrinkage estimator:

$$\hat{s}_g^2 = \frac{(IJ-1)s_g^2 + d\hat{s}_g^2}{IJ-1+d}, \quad (17)$$

where  $d$  is the degree of freedom of  $\hat{s}_g^2$ . The estimator (17) is the Bayes estimator with the inverse Gamma prior with the prior mean

$\hat{s}_g^2$  and prior degree of freedom  $d$ . Since the prior parameters are estimated from the data, (17) is indeed an empirical Bayes estimator.

The degree of freedom of the nonparametric estimator  $\hat{s}_g^2$  is often estimated by the effective sample size, which is the inverse of the asymptotic variance of the kernel local linear regression estimator. For example, if the kernel local regression estimator is used, then, we have

$$d = \text{the number of local data} / (2\|K\|^2),$$

where  $K$  is the kernel function used in the smoothing. For example, in the LOWESS smoother, the kernel function is  $\frac{70}{81}(1 - |x|^3)^3 I(|x| \leq 1)$ .

Note that when the within-array variance  $s_{W,g}^2$  in (13) is used in (17), the factor  $(IJ - 1)$  should be replaced by  $I(J - 1)$ . In addition, the intensity-specific estimate of variance is usually not as reliable as the within-array variance  $s_{W,g}^2$ , the constant  $d$  can be replaced by some smaller numbers to reduce somewhat of its influence.

## 5 SIMULATIONS AND APPLICATIONS

In this section, we first use the simulated data set to illustrate the validity and the power of our proposed validation tests. In particular, the advantages of the aggregated standardization tests  $T_3$  and  $T_4$  are demonstrated. The three real data analyses illustrate the methodological power of our approaches.

### 5.1 Simulation

In each simulation, we generate  $J = 4$  arrays from model (3) with  $G = N = 2000$  genes, each having  $I = 3$  replications randomly assigned over the 48 blocks. The details of simulation scheme for this example are summarized as follows:

$\alpha_g$ : The expression levels of the first 150 genes are generated from the standard double exponential distribution. The rest are 0's. These expression levels are the same over 4 arrays in each simulation, but may vary over simulations.

$\beta$ : The 48 parameters for the block effects  $\beta_j$  in array  $j$  are all set to  $\beta$ , which is given by

$$\begin{aligned} \beta = & (-0.15 \ -0.3 \ -0.15 \ 0.01 \ -0.01 \ 0.04 \ 0.07 \ 0.08 \ 0.08 \ 0.23 \\ & 0.22 \ 0.09 \ 0.11 \ 0.24 \ 0.28 \ 0.24 \ 0.07 \ 0.22 \ 0.19 \ 0.15 \\ & -0.03 \ -0.05 \ -0.02 \ 0.03 \ -0.14 \ -0.04 \ -0.18 \ 0.06 \ -0.05 \ -0.02 \\ & -0.17 \ -0.02 \ 0.08 \ 0.08 \ -0.04 \ -0.16 \ 0.04 \ 0.01 \ -0.05 \ -0.12 \\ & 0.07 \ -0.19 \ -0.03 \ -0.15 \ -0.07 \ -0.22 \ -0.11 \ -0.22)^T. \end{aligned}$$

These parameter values are taken from the estimates in Ma *et al.*, 2006.

$X$ : The intensity is generated from a mixture distribution: with probability 0.8 from probability distribution  $.0004(16 - x)^3 I(6 < x < 16)$  and 0.2 from the uniform distribution over  $[4, 16]$ .

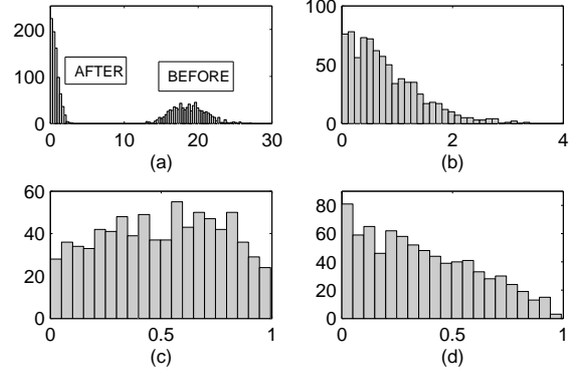
$m_j(\cdot)$ : Set the function  $m_j(X) = 5(.3592 - \sqrt{(X - 4)/32})$ , whose expectation with respect to  $X$  is approximately zero.

$b_g$ : For each given gene, its associated block is assigned at random at one of 48 print-tip blocks.

$\varepsilon$ :  $\varepsilon_{gij}$  is generated from the standard normal distribution with mean zero and variance  $\sigma^2(X_{gi}) = .15 + .015(X_{gi} - 9)^2 I\{X_{gi} > 9\}$ .

This is a heteroscedastic model with small block effect and intensity effect. Three normalization methods are used: Global

normalization, LOWESS normalization using all blocks, and the SLIM normalization (Fan *et al.*, 2004). They are applied to 200 pseudo-data sets, each having  $J = 4$  arrays, giving a total of 800 arrays. We drew 200 genes at random from the 2,000 different genes as the testing set, and the remaining genes as the learning set.

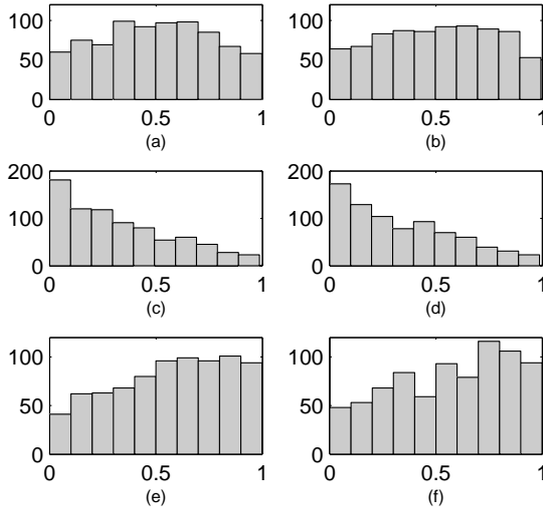


**Fig. 1.** (a): Distributions of  $|T_3|$  before normalization (right) and after normalization using SLIM (left). (b): Distribution of  $|T_3|$  after the LOWESS normalization without accounting small block effects. (c) and (d): Distributions of P-values of the validation test  $T_3$  after normalization using SLIM (left panel) and LOWESS (right panel) methods. P-values based on test  $T_3$  before normalization are all zero.

We first apply the validation test statistic  $T_3$  with genewise variances estimated by the REML estimators (14) to check the effectiveness of the three normalization methods over 200 pseudo-data sets, which comprise of 800 pseudo-arrays. The distributions of the test statistic  $|T_3|$  based on the three normalization methods are presented in Figure 1. They are well separated (Fig. 1(a)). First of all,  $T_3$  for the global normalization is the same that with no normalization, which by far the largest. This shows the global normalization is the worst method for the data sets. Indeed, the corresponding 800 P-values are all zero, which shows the power of validation test is 100%. The LOWESS normalization using all blocks ignores the small block effects  $\beta$ . The resulting statistics  $T_3$  from 200 simulations are smaller than those based on the global normalization, but are stochastically larger than those based on the SLIM normalization, which accounts for these small block effects. This shows that the LOWESS normalization ignoring block effect is not as effective as the SLIM, but is more effective than the global normalization. Figure 1 also depicts the distribution of the 800 P-values of the LOWESS and SLIM method. The P-values of SLIM follow nearly a uniform distribution, which indicates that systematic biases have been effectively removed. On the other hand, for a portion of arrays, the LOWESS normalization is inadequate. This demonstrates the power of the validation: even ignoring small block effects, the test statistic  $T_3$  is able to detect these small systematic biases.

After demonstrating the power of the test statistic  $T_3$ , we now examine the accuracy of the null distributions  $T_1, \dots, T_4$  using SLIM as the method of normalization. Since the correct method is used in the normalization for the pseudo-data, the estimation errors come from two sources: estimation of block and intensity effect and estimation of genewise variance by (14). First of all,

without normalization, all validation test statistics  $T_1, \dots, T_4$  report zeros P-values for all realizations. This indicates the sensitivity of these tests. When the SLIM is used to normalize the data, if the normalization method is effective and the null distributions are accurate, the P-values follow the uniform distribution on the interval  $[0, 1]$ . Figure 2 depicts the P-values based on the SLIM normalization. The distributions of P-values based on  $T_3$  are reasonably uniform. This shows that both the normalization method



**Fig. 2.** Distributions of P-values for test statistics  $T_1, T_2, T_3$ , and  $T_4$  after the SLIM normalization. (a)-(d): Distributions of P-values for test statistics  $T_1, T_3, T_2$ , and  $T_4$  after the SLIM normalization. (e) and (f): Distributions of P-values for test statistics  $T_2$  and  $T_4$  calculated by using unbiased estimators of genewise standard deviation (12). The distribution of  $T_3$  is somewhat more uniform than that based on  $T_1$ . The same conclusion applies to  $T_4$  and  $T_2$ , both using the REML estimators and unbiased REML estimators.

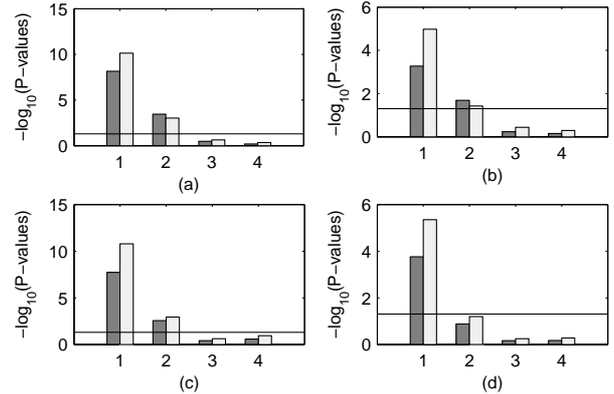
is effective and the null distribution is accurate for the test statistic  $T_3$ . However, the distributions of  $T_2$  and  $T_4$  have large deviations from the uniform distribution, which indicates the estimation errors from the REML estimators (14) of genewise standard deviations. These deviations have greatly been mitigated by correcting the REML estimators to the unbiased estimators (12).

### 5.2 Application to human placenta data

A collection of human placenta cDNAs comprising 7042 clones was identified and used as the probe set for cDNA microarray fabrication in this study (Ma *et al.*, 2005). Three kinds of RNA samples were used. These include the common reference RNA derived from the probe set (PS) in equal amounts representing artificial RNA produced by in vitro transcription, the ‘‘Universal Human Reference RNA’’ from Stratagene, which is comprised of 10 different cell lines and human full-term placenta RNA. The original goal of the study was to evaluate the performance of the PS RNA as a reference RNA in comparison with that of Stratagene’s universal reference RNA.

For the sake of studying the normalization and validation tests, we only compare the ‘‘Universal Human Reference’’ RNA with human placenta RNA in this study. Gene expression values were obtained through direct hybridizations between these two kinds of

RNAs. There are four slides, including two dye-swapped slides. Each clone was printed three times on different blocks in each slide. There are 48 blocks on each array. After preprocessing that filters low quality spots, there remained 2149 genes that have 3 replications. Our analysis focuses on this subset. We compared the effectiveness of four normalization methods: Global, LOWESS, SLIM and aggregated SLIM by using test statistics  $T_3$  and  $T_4$ . One hundred different genes were selected as the validation set, while the remaining 2049 genes were used to estimate the parameters.



**Fig. 3.** The P-values for validation tests  $T_3$  and  $T_4$ , based on the human placenta data, under four different normalization approaches: Global (labeled ‘‘1’’), LOWESS (labeled ‘‘2’’), SLIM (labeled ‘‘3’’), and aggregated SLIM (labeled ‘‘4’’). The y-axis is  $-\log_{10}(\text{P-value})$ . The dark grey columns are P-values for  $T_3$ , while the light grey ones are for  $T_4$ . The results for four arrays are plotted in (a) – (d). The lines correspond to 5% significance level. The P-values for Global normalization are highly statistically significant at level 5% for all 4 arrays, indicating the ineffectiveness of the method. The LOWESS normalization method improves the P-values a lot but still are statistically significant at level 5% except the 4<sup>th</sup> array. The P-values based on SLIM and aggregated SLIM methods show that the systematic biases due to the variation of experimental conditions have been effectively removed by either of these two methods. Therefore, the resulting log-ratios can be used for the downstream statistical analysis.

The key assumption of the LOWESS method is that up- and down-regulated genes’ expressions are symmetrically distributed around 0, which is usually not the case for genes investigated in placenta tissue (Ma *et al.*, 2005). Figure 3 (a)-(d) compares the effectiveness of the four normalization methods using the validation test statistics  $T_3$  and  $T_4$ . The genewise variances are estimated by (14). The results show clearly that normalizations are needed for each array. In addition, the blockwise LOWESS normalization is inadequate except the 4<sup>th</sup> array at significant level 5%. The outcomes of validation tests show that we can choose either SLIM or aggregated SLIM to normalize the log-ratios for each array.

### 5.3 Application to interferons data using customized arrays

An important property of interferons (IFNs), a cytokine, is their anti-tumor activity. IFNs have efficacy in the treatment of several types of solid tumors/carcinomas. Interestingly, it has been reported that IFN- $\beta$  has greater anti-tumor effects than IFN- $\alpha$  on melanoma. One

probable mechanism for the different effects may be the different affinity of IFN- $\alpha$  and IFN- $\beta$  binding to the IFN receptors. Another possibility may be the differences in intracellular signaling. To address whether different signal pathways are involved in IFN- $\alpha$  and IFN- $\beta$ -mediated anti-tumor activity, customized c-DNA chips are designed to include IFN stimulated genes and genes involved in multiple pathways. Gene expression changes induced by IFN- $\alpha$  and IFN- $\beta$  were investigated and compared by using the customized c-DNA microarrays.

The customized c-DNA microarrays provides an ideal platform for our validation tests. Usually, only several hundreds of genes of primary interest are monitored for the changes of expression profiles and these genes are often duplicated several times. For our particular applications, 768 genes that might be induced by IFN are printed on the 16 blocks with 12 rows and 12 columns of spots in each block. These 768 genes are duplicated 3 times. The three replications of each gene reside in the same row in the same block but adjacent columns. The expression profiles of these 768 genes with IFN- $\alpha$  or IFN- $\beta$  stimulations are compared with those without stimulations. This results in two customized microarrays, one with INF- $\alpha$  stimulation and the other with INF- $\beta$  stimulation, each having with  $3 \times 768$  spots. After some preprocessing that filtered the data with low quality, there were 572 genes left for further analysis.

To examine the necessity and effectiveness of normalization, we randomly selected 50 genes as the test set; the remaining 522 genes were used as the learning set. Three normalization methods are employed: Global, LOWESS and SLIM normalizations. Due to the specific designs of the replications, the block effect is not estimable. The genewise variance are estimated by using the empirical Bayes method in section 4.3 – the variance estimates from two arrays are not aggregated. Table 2 depicts the P-values using the test statistics  $T_1, \dots, T_4$ . From the table, we conclude that there is no need for normalization since all of the P-values for test statistics are high for both slides and for all methods. Note that the P-values by using the SLIM normalization are a little bit higher than those using the Global normalization, and they are larger than the P-values using the LOWESS method. That is because the fundamental assumption for LOWESS method – up- and down- regulated genes are about the same – are not substantiated here.

Table 1. P-values for  $T_1, \dots, T_4$  based on human placenta data

		P-values		
		Global	LOWESS	SLIM
Slide 1	$T_1$	0.3294	0.2857	0.3333
	$T_2$	0.3979	0.3139	0.4044
	$T_3$	0.5431	0.5155	0.5450
	$T_4$	0.3947	0.3097	0.4004
Slide 2	$T_1$	0.4704	0.4966	0.4730
	$T_2$	0.6952	0.6615	0.6964
	$T_3$	0.3012	0.2261	0.3009
	$T_4$	0.4322	0.3913	0.4318

### 5.4 Application to human total RNA samples using Agilent arrays

Our third example comes from the Microarray Quality Control (MAQC) project (Patterson *et al.*2006). The MAQC project studies

the reproducibility, sensitivity and specificity of the microarray data across different platforms and sites. It compared two RNA samples, Stratagene Universal Human Reference total RNA and Ambion Human Brain Reference total RNA using different microarray technology. Our study focuses only on the RNA samples generated at three test sites using Agilent platform. At each site, ten Agilent two-color microarrays were processed with five arrays for each dye configuration, which assayed a total of 30 microarrays. Following Patterson *et al.*(2006), we excluded two outlier microarrays based on single microarray quality metrics, resulting in 28 microarrays. After preprocessing, we obtained 21,767 genes from a total of 43,931 and found four genes each having 10 replications, randomly located on the microarray.

For our validation test, gProcessedSignal and rProcessedSignal values from Agilent’s Feature Extraction software were used as input to calculate the test statistics. If the genewise variance is estimated by using (14), all P-values are at least 99.99%, indicating the proper normalization of all Agilent arrays. Even using the most stringent estimation of genewise variance (13), almost all the microarrays pass the validation test at significant level 1% except for one array AGL-3-D3 at the test site 3. See Table 2. The results show the data processed by Agilent software are properly normalized and reliable. These are in line with the conclusion of the MAQC project.

Table 2. P-values based on  $T_1, \dots, T_4$  for MAQC project data

slide name	P-values			
	$T_1$	$T_2$	$T_3$	$T_4$
AGL-1-C1	1.0000	1.0000	0.9993	0.9999
AGL-1-C2	1.0000	1.0000	0.9989	0.9996
AGL-1-C3	1.0000	1.0000	0.9982	0.9997
AGL-1-C4	1.0000	1.0000	0.9992	1.0000
AGL-1-C5	0.9998	1.0000	0.9962	0.9957
AGL-1-D2	0.2706	0.9948	0.2965	0.3792
AGL-1-D3	0.6940	0.9993	0.6931	0.7336
AGL-1-D4	0.5144	0.9981	0.5334	0.5785
AGL-1-D5	0.9894	1.0000	0.9659	0.9993
AGL-2-C1	0.7041	0.9977	0.6903	0.5178
AGL-2-C2	0.9479	0.9999	0.9247	0.9396
AGL-2-C3	0.9794	1.0000	0.9632	0.9919
AGL-2-C4	0.4056	0.9932	0.3823	0.2779
AGL-2-D1	0.0848	0.9502	0.0821	0.0562
AGL-2-D2	0.2218	0.9688	0.2497	0.1067
AGL-2-D3	0.3557	0.9889	0.3980	0.2688
AGL-2-D4	0.0177	0.6857	0.0109	0.0004
AGL-2-D5	0.1242	0.9551	0.1272	0.0662
AGL-3-C1	0.0570	0.8793	0.0451	0.0085
AGL-3-C2	0.0389	0.7861	0.0291	0.0019
AGL-3-C3	0.1024	0.9474	0.1033	0.0505
AGL-3-C4	0.1203	0.9092	0.1276	0.0215
AGL-3-C5	0.1328	0.9393	0.1337	0.0366
AGL-3-D1	0.2748	0.9645	0.3138	0.0888
AGL-3-D2	0.1372	0.8813	0.1371	0.0094
AGL-3-D3	<b>0.0034</b>	0.5220	<b>0.0007</b>	<b>0.0000</b>
AGL-3-D4	0.1907	0.9694	0.2071	0.1004
AGL-3-D5	0.0388	0.8792	0.0348	0.0115

## 6 CONCLUSION

Motivated by the urge of measures for comparing different normalization methods, we proposed four validation tests to evaluate the necessity and effectiveness of normalization methods, relying on the replicated clones. They are based on the standardized differences of expression profiles among replicated clones aggregated over different genes, resulting in the test statistics  $T_1$  and  $T_2$ . These tests depend on genewise variances and standard deviations and hence can not be used to compare the effectiveness of the normalization without estimation of these genewise variances. This leads us to consider the (unweighted) differences of expression profiles among replicated clones aggregated over different genes, standardized after aggregation, resulting in the test statistics  $T_3$  and  $T_4$ . The unscaled test statistics  $T_3$  and  $T_4$  can be used to compare the effectiveness of normalization without estimating genewise variance (which itself depends on selecting a method of normalization). The aggregated standardization tests  $T_3$  and  $T_4$  depend on the aggregated SD and variance, which can be more precisely estimated than the genewise standard deviation and variance. As a result, the null distribution of the test statistics  $T_3$  and  $T_4$  can be more accurately approximated.

We have also demonstrated that the within-array replications are essential for estimating the genewise variance. A novel nonparametric approach is proposed, which aggregates variance information from genes with similar intensity level. Several innovative approaches are proposed to enhance the accuracy of the estimation of the genewise variance. These new methods can also be used to improve the power of selecting significantly differently expressed genes (Cui *et al.*, 2005; Smyth *et al.*, 2005).

Our simulation studies show convincingly the power of the validation tests and their validity. The applications to three real data sets demonstrate the methodological power of our proprietary methods in choosing a normalization method and in assessing whether the systematic biases due to variations in the experimental conditions have been properly removed. The customized array provides an ideal platform for the applications of our proposed approaches. Furthermore, our idea doesn't restrain to the two-color arrays. As long as replicated genes exist, we could apply our validation test to check the necessity and effectiveness of normalization. The validation tests could have been reliably applied the one-color Affymetrix GeneChip arrays, if they were within-array replications. Without such replications, we need to aggregate information across arrays. Assuming the absence of the gene and array interactions, our validation tests can be applied to the Affymetrix array to check the the necessity and effectiveness of normalization, by treating each array as a block in a synthetic super-array.

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