

Physically grounded approach for estimating gene expression from microarray data

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High-throughput technologies, including gene-expression microarrays, hold great promise for the systems-level study of biological processes. Yet, challenges remain in comparing microarray data from different sources and extracting information about low-abundance transcripts. We demonstrate that these difficulties arise from limitations in the modeling of the data. We propose a physically motivated approach for estimating gene-expression levels from microarray data, an approach neglected in the microarray literature. We separately model the noises specific to sample amplification, hybridization, and fluorescence detection, combining these into a parsimonious description of the variability sources in a microarray experiment. We find that our model produces estimates of gene expression that are reproducible and unbiased. While the details of our model are specific to gene-expression microarrays, we argue that the physically grounded modeling approach we pursue is broadly applicable to other molecular biology technologies.

process modeling | statistical power

One thousand manuscripts are published each year involving microarray technology.† In spite of the 15-year history of the field, those manuscripts still describe a wide variety of data analysis methods, many of them poorly specified. Indeed, criticisms of the validity and reproducibility of microarray experiments have dogged the technology since its inception. There are two possible explanations for these shortcomings: (i) inherent limitations of the microarray technology that constrain its utility or (ii) modeling strategies that are not appropriate. The former is potentially a fundamental problem that can be overcome only with technological advances. This hypothesis has led to candid speculation that emerging sequencing technologies will quickly replace microarrays as the de facto genome-wide expression analysis technique (1, 2).

An alternative view is that current shortcomings result from gaps in our understanding of how to model the data generated in microarray experiments. In order to pursue this point, let us consider the motivation for the “standard” model (3). The fluorescence intensity F_i (Fig. 1A) detected at a spot i is surmised to be the sum of a background term and a term related to the expression level E_i we want to estimate,

$$F_i = B_i + f(E_i). \quad [1]$$

Oddly, the standard model assumes that B_i can be directly determined from the fluorescence intensity measured in the nonfeature region surrounding the spot.‡ The dependence on E_i is assumed to be distorted by multiplicative noise (3). These assumptions yield

$$F_i = B_i^{\text{nf}} + E_i A_i e^{\nu_i^{\text{sp}}}, \quad [2]$$

where ν^{sp} is normally distributed with zero mean, and A_i is a parameter capturing the effects of hybridization efficiency and dye-specific and experiment-specific factors.

Because of the difficulty in estimating systematic effects affecting the value of A , microarray experiments are frequently performed with an internal control, the goal being to determine

change of expression R_i between two conditions, 1 and 2, instead of the expression level for each condition:

$$\hat{R}_i = \log\left(\frac{\hat{E}_i^1}{\hat{E}_i^2}\right) = \log\left(\frac{F_i^1 - B_i^1}{F_i^2 - B_i^2}\right) + A_i', \quad [3]$$

where $A_i' = \log(A_i^2/A_i^1)$, \hat{R}_i and \hat{E}_i are the best estimates of R_i and E_i . Because, according to Eq. 2, F_i and B_i can be directly measured, the crux of the traditional approach is to estimate A_i' .

In dye-swap experiments, for which the two conditions are identical, one can develop a number of reasonable expectations for $p(R_i)$ and $p(R_i|E_i)$. Assuming no correlations in the values of A_i' , one expects the average value of R_i to be zero. Moreover, assuming that A_i is nonnegligible, one expects the standard deviation of R_i to decrease with increasing E_i . Unfortunately, neither of these expectations is typically obeyed by the data (Fig. 1 B, C, D).

As a result, the field has failed to converge on a single, robust model. Instead, publications reporting microarray data include a bevy of variations of this standard model. In many cases, these models were “rescued” to achieve the aforementioned expected properties by the use of idiosyncratic nonlinear corrections. Exemplifying this are the data reanalyzed in this manuscript—the authors of the studies considered have used different normalization techniques (4, 5).

Here, we argue that background fluorescence intensity cannot be correctly estimated by B_i^{nf} . Nonspecific hybridization is the dominant factor determining B_i . In order to correctly estimate B_i , we propose a dramatically distinct approach to determining gene-expression levels from microarray data. Instead of attempting to surmise a functional expression for F_i , we model each of the processes that constitute a microarray experiment. Remarkably, by propagating the fluctuations one expects in each stage of the protocol, we arrive at a concise expression relating E_i to measured quantities in the experiment.

We find that our model is able to capture the properties of microarray data for thousands of experiments. Moreover, our model yields reproducible estimates of changes in expression level.

The Physically Grounded Approach

The protocol for two-color cDNA microarray experiments is now essentially standard (6). The measurement component has three

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‡Due to differences in surface chemistry of feature and nonfeature regions, one cannot reasonably expect that B_i is representative of the background fluorescence in the feature region.

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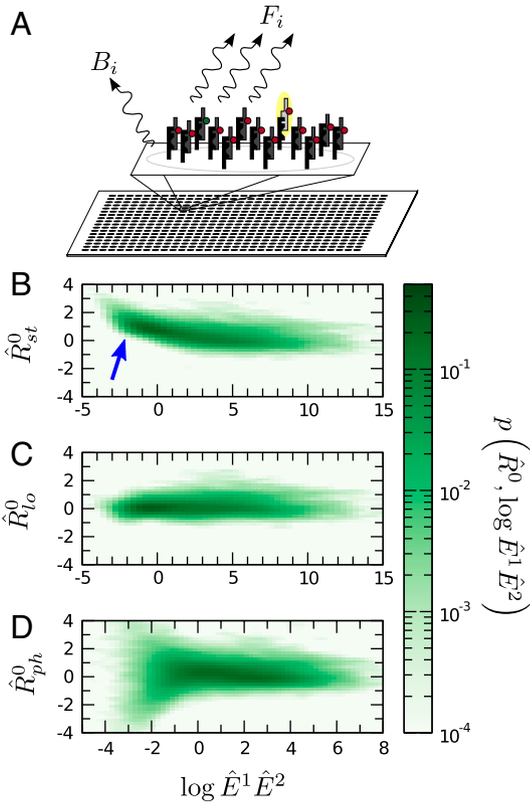


Fig. 1. (A) Schematic of a microarray chip. Two quantities are typically reported for each spot in a two-color microarray experiment: the median feature fluorescence F_i and the median nonfeature fluorescence B_i . The total feature intensity is constituted by the sum of intended specific associations between probe and target (dark gray), as well as any number of nonspecific interactions (light gray). Because the nonfeature region has no probes attached, it is unreasonable to assume that B_i can provide reliable information on the nonspecific hybridization occurring in the feature region. (B–D) Single chip-wide joint probability distributions of \hat{R}^0 and $\log \hat{E}^1 \hat{E}^2$ for an *A. thaliana* chip (19) (GEO accession no. GSM133484). (B) A plot of \hat{R}^0 against $\log \hat{E}^1 \hat{E}^2$ is equivalent to the “MA” plots commonly used to diagnose bias in microarray data. \hat{R}^0 estimates from the statistical model (Eq. 2) depend strongly on \hat{E} , particularly for small \hat{E} (blue arrow). This bias is absent from data adjusted using (C) the scatterplot smoothing routine lowess and (D) from estimates derived from our physically grounded model.

main stages, each with its own characteristics. Sample preparation consists of mRNA extraction, purification, amplification, and labeling. Hybridization is the process by which differently labeled targets bind surface-associated probes. Detection is the excitation and scanning of surface-associated fluorophores. In the following, we describe and model each of these stages.

Consider a biological sample consisting of E_i copies of transcript i , with $i = 1, \dots, N_{\text{tr}}$. The quantity of RNA derived from a biological sample is typically insufficient for efficient quantification by current experimental methods. Thus, sample amplification is necessary. One of two methods is typically employed to amplify the original messenger RNA: (i) expression in a T7 viral vector or (ii) polymerase chain reaction. Amplification by T7 vector expression is currently the preferred method because it results in smaller variability for high expression levels (7); thus, we consider it here (SI Text).

cDNA vectors are prepared from sample mRNAs by incorporating the T7 polymerase promoter into reverse transcriptase primers. Approximately one vector arises from each mRNA. We assume that transcription of these vectors to RNA is kinetically limited by the rate R_b of binding of T7 polymerase to transcription start sites (8). In our model, we disregard sequence or length dependent effects on transcription rate (7, 9).

In a well-mixed solution, transcripts of gene i are produced at a characteristic rate, $E_i R_b$. Under experimental conditions, the number of transcripts present after running the process for a time t is described by a Poisson process with parameter $E_i R_b t$. We expect the amplification gain to be very high, that is, $R_b t \gg 1$. In this limit, the Poisson distribution of number of transcripts arising from this process converges to a Gaussian distribution. This implies that the number n_i of copies of cDNA for gene i available for hybridization is a Gaussian variate with mean and variance equal to $E_i R_b t$.

Consider now competitive hybridization in a solution that is well-mixed and let p_{ii} be the probability of specific hybridization of target i to feature i . p_{ii} may depend on the sequence of gene i and on experimental conditions such as temperature and buffer concentration, but typically probe sequences are selected so that p_{ii} is approximately constant. Thus, we assume that $p_{ii} = p_{\text{sp}}$ for all i and let its fluctuations be incorporated into the noise. We suggest that the number S_i^{sp} of specifically hybridized probes in the feature follows a binomial distribution with parameter p_{sp} . If $n_i p_{\text{sp}} \gg 1$, then the central limit theorem holds, and S_i^{sp} is a Gaussian variate with mean $n_i p_{\text{sp}}$,

$$S_i^{\text{sp}} = E_i R_b t p_{\text{sp}} (1 + \epsilon_i^t) (1 + \epsilon_i^h), \quad [4]$$

where ϵ_i^t and ϵ_i^h are Gaussian variates with zero mean.

Similarly, let p_{ji} be the nonspecific hybridization efficiency for gene j to probe i . The number of hybridized probes j in feature i will then be

$$S_{ji} = n_j p_{ji} (1 + \epsilon_{ji}^h), \quad [5]$$

where ϵ_{ji}^h is again a Gaussian variate with mean zero. Note that $p_{ji} \ll p_{ii}$ for all $j \neq i$. The total contribution of nonspecific hybridization from all targets to the observed signal will then be

$$S_i^{\text{ns}} = \sum_{j \neq i} S_{ji} = \sum_{j \neq i} [n_j p_{ji} (1 + \epsilon_{ji}^h)]. \quad [6]$$

Estimating p_{ji} directly for all pairs of transcripts is not feasible in practice. In order to proceed, we thus use a mean-field approximation. Specifically, we assume that no single gene is responsible for a significant fraction of all mRNA targets. We further assume that p_{ji} is not dependent strongly on j or i ; that is $p_{ji} \approx p^{\text{ns}}$. Under these assumptions, Lyapunov’s central limit theorem applies, yielding

$$S_i^{\text{ns}} = U' (1 + \epsilon_i^{\text{ns}}), \quad [7]$$

where U' is the characteristic contribution of nonspecific hybridization and ϵ_i^{ns} is a Gaussian variate with zero mean.

The fluorescence generated by the excitation of the spots on the chip will be amplified in the scanning process. Amplification using a photomultiplier is characterized by a dye-specific gain G that is a function, in principle, of dye incorporation rate, dye properties, laser power, and detector characteristics, yielding a detected fluorescence

$$F_i = (S_i^{\text{ns}} + S_i^{\text{sp}}) G_i \prod_{k=1}^m (1 + \epsilon_i^k), \quad [8]$$

where ϵ_i^k , the noise associated with stage k of amplification, is normally distributed with mean zero. We assume that the gain is constant and does not depend on the intensity of the signal, or on any other spot property; that is, $G_i = G$. We also assume that there is no specific interaction between a dye molecule and either target or probe sequence, an assumption that we find fails for some probes (SI Text).

Because the variability for each term in Eq. 8 is the product of several independent Gaussian variables, the terms will converge to a log-normal distribution. We can therefore write Eq. 8 as

that is unobservable using other models. *This leads us to conclude that our model imparts greater statistical power.*

Having established the statistical legitimacy of our model, we investigated what practical implications it has for identifying consistent sets of up- and down-regulated transcripts. For our model and the three others, we determined the 100 genes most likely to be up- and down-regulated for each chip. For the HC and HCR chips, we aggregated genes that were represented in at least three of four sets (Fig. 6C). Given the hypothesis that HCR animals are similar to control animals, we expect that there is much less consistency between the HCR sets (i.e., small circles). Also, if the HCR and HC animals are distinct, they should have very few genes common across conditions (i.e., small overlap of the circles). This is the behavior we observe in the sets derived from our model—there is very little consistency between the HCR sets, but the HC sets are robust—but *not* for the other models.

Discussion

We demonstrate here that a physically grounded approach successfully models the outcome of gene-expression microarray experiments. Whereas linear models assuming normally distributed error terms may be appropriate models for many experiments in biology, they fail in many high-throughput applications due to the multiplicative nature of propagating fluctuations. For these experiments, consideration of the physical processes responsible for the outcome is essential. Our model, although constructed with two-color expression microarrays in mind, is generalizable to other systems. As chip-based assays and other high-throughput technologies continue to evolve, it will become increasingly important to establish physically grounded models for the resulting data. Although the specifics of a particular protocol may vary, a physically grounded model can be derived to understand any procedure that is composed of serial, fundamentally understood

stages. For these experiments, statistical models are often the first approach because physically grounded models may be perceived as difficult to develop. In many cases, the benefits of the physically grounded modeling approach are appreciable and may outweigh increased developmental difficulty.

We have found that this approach produces a model for microarray data that reproduces macroscopic properties of the chip and results in estimates of expression changes that are nearly free of intensity-dependent dye bias, an artifact that has been traditionally rectified using ad hoc approaches. As a result, the estimates we obtain of the expression levels are systematically reproducible within and across laboratories. In addition, our model allows us to assign confidence to expression changes, even in experiments devoid of technical and biological replicates.

Our study provides yet another cautionary tale of the ad hoc adjustment of models of complex data. The standard statistical model of microarray data has many laudable features: It is simple, it has easily understood parameters, and it is readily testable. Indeed, the model's inability to capture even basic properties of the data (Figs. 1, 2, and 3) would strongly suggest the need to reject it. Surprisingly, instead of rejecting the model, the course followed by the field has been its "rescuing" with uncontrolled and unjustified corrections. Our study shows that these corrections are unnecessary.

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