Empirical Bayes Analysis of a Microarray Experiment

Bradley Efron, Robert Tibshirani, John D. Storey, and Virginia Tusher

Microarrays are a novel technology that facilitates the simultaneous measurement of thousands of gene expression levels. A typical microarray experiment can produce millions of data points, raising serious problems of data reduction, and simultaneous inference. We consider one such experiment in which oligonucleotide arrays were employed to assess the genetic effects of ionizing radiation on seven thousand human genes. A simple nonparametric empirical Bayes model is introduced, which is used to guide the efficient reduction of the data to a single summary statistic per gene, and also to make simultaneous inferences concerning which genes were affected by the radiation. Although our focus is on one specific experiment, the proposed methods can be applied quite generally. The empirical Bayes inferences are closely related to the frequentist false discovery rate (FDR) criterion.

1. INTRODUCTION

Through the use of DNA microarrays, a novel technology, it is now possible to obtain quantitative measurements for the expression of thousands of genes present in a biological sample. DNA microarrays have been used to monitor changes in gene expression during important biological processes (e.g., cellular replication and the response to changes in the environment), and to study variation in gene expression across collections of related samples (e.g., tumor samples from patients with cancer). A major statistical task is to understand the structure of the data from such studies, which often consist of measurements on thousands of genes in dozens of conditions.

This article concerns the use of microarrays in a comparative experiment, where it is desired to compare gene expression under Treatment versus Control conditions. We wish to identify which of several thousand candidate genes have had their expression levels changed, either positively or negatively, by the Treatment. Answering this question requires an efficient data reduction strategy, because microarrays deliver megabytes of information, and also statistical inference techniques that deal with the difficulties of simultaneous inference on thousands of genes. We discuss both problems here, working in the context of an experiment on radiation sensitivity discussed later.

The statistics literature for microarrays, still in its infancy and with much of it unpublished, has tended to focus on frequentist data-analytic devices such as cluster analysis, bootstrapping, and linear models (see Li and Wong 2000; Kerr and Churchill 2000; Black and Doerge 2000; Van del Laan and Bryan 2000; Eisen, Spellman, Brown, and Botslein 1998). Parametric Bayesian modeling was featured in Newton, Kendziorski, Richmond, Blatter, and Tsui (2000) and to a less extent in Lee, Kuo, Whitmore, and Sklar (2000). Multiple comparison techniques, designed to control error rates in thousands of simultaneous hypotheses tests, were explored in Dudoit, Yang, Callow, and Speed (2000). Tusher, Tibshirani,

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and Chu (2001) approached the simultaneity problem through the method of false discovery rates (FDRs), as discussed later.

Our inferences here will be based on a simple nonparametric empirical Bayes model. The model produces useful *a posteriori* probabilities of effect for the individual genes, with a minimum of prior assumptions. It also connects well with Benjamini and Hochberg's (1995) frequentist theory of FDRs, as discussed in Section 5. Besides being useful in its own right, the empirical Bayes model helps to select from among competing data reduction schemes, a crucial point in dealing with the massive datasets microarrays produce.

Here is some background on microarrays in general and the specific experiment analyzed in this article. Virtually all living cells contain chromosomes, large pieces of DNA containing hundreds or thousands of genes, each of which specifies the composition and structure of a protein (or sometimes several related proteins). Protein polymers of amino acids are the workhorse molecules of the cell, responsible, for example, for cellular structure, producing energy and important biomolecules like DNA and proteins, and for reproducing the cell chromosomes. Every cell in an organism has nearly the same set of chromosomes, and thus contains the same repertoire of proteins. However, cells have remarkably distinct properties, such as the differences between human eye cells, hair cells and liver cells, distinctions that are the result of differences in the abundance, distribution, and state of the cell proteins. One of the seminal discoveries of molecular biology was that these changes in protein abundance are determined in part by changes in the levels of messenger RNA (mRNA), small and relatively unstable nucleic acid polymers that shuttle information from chromosomes to the cellular machines that synthesize new proteins. Thus, there is a logical connection between the state of a cell and the details of its protein and mRNA composition.

Whereas it remains difficult to measure the abundances of a cell's proteins, the recently developed DNA microarray makes it possible to quickly and efficiently measure the relative representation of each mRNA species in the total cellular mRNA population, or in more familiar terms to measure gene expression levels.

There are two major kinds of microarrays. In an oligonucleotide array, the kind featured in this article, there are 20 probe pairs (pm, mm) for each gene. The perfect match (pm)

probe is designed to match a small subsequence of the gene about 25 bases long. The mismatch (mm) probe is a control, being identical to pm except with the middle base flipped to its complement. An experimental sample is hybridized on the microarray, and the RNA expression of the gene is estimated by the difference in signal pm—mm averaged over the 20 probe pairs. There is some concern that subtracting mismatch numbers may actually degrade the inferences, a question we consider in this article.

In a spotted cDNA microarray, the other major variety, one base sequence matching all or part of a gene is printed on a glass slide. The experimental sample is labeled with red dye and hybridized on the slide. As a control, a reference sample is labeled with green dye and hybridized on the same slide. Using a fluorescent microscope, the log (red-green) intensities of RNA hybridization at each site are measured. The redgreen microarray is featured in much of the recent literature, see Newton et al. (2001), Dudoit et al. (2000), and Lee et al. (2001). Our discussion, like that in Li and Wong (2000) centers in the Affymetrix oligonucleotide microarray, but similar analysis problems arise for both types of array. However, our Empirical Bayes procedure, summarized in Algorithm 1, can be applied quite generally. An example extending the empirical Bayes analysis to a cDNA microarray experiment appears in Remark D of Section 6, showing how our methods can be applied to other experimental situations.

From either type of microarray, we obtain several thousand expression values, one or many for each gene. Microarrays in current use measure anywhere from 1,000 to 25,000 genes; larger ones will soon be available. In a typical study, a number of experimental samples are each hybridized to a different microarray to learn about gene expression differences across different conditions. For example Alizadeh et al. (2000) studied gene expression patterns from tissue samples from a number of lymphoma patients and related gene expression to patient survival. Clustering methods (Eisen et al. 1998) were the main tool used in that article, and in a number of other similar studies. Here we will be interested in the more familiar statistical task of comparing Treatment and Control arrays, though carried out in a novel setting.

Our particular dataset comes from a set of eight oligonucleotide microarrays in an experiment designed by Professor Gilbert Chu of the Stanford Biochemistry Department to study transcriptional responses to ionizing radiation. Some cancer patents have severe life-threatening reactions to radiation treatment. It is important to understand the genetic basis of this sensitivity, so that such patients can be identified before the treatment is given. The eight microarrays were labeled

$$(U1A, U1B, I1A, I1B, U2A, U2B, I2A, I2B),$$
 (1.1)

the labels indicating the following experimental design: RNA was harvested from two wild-type human lymphoblastoid cell lines, designated "1" and "2," growing in an unirradiated state "U," or in an irradiated state "I." RNA samples were labeled and divided into two identical aliquots for independent hybridizations, "A" and "B." Each microarray provided expression estimates for 6,810 genes. Further experimental details appear in Remark A of Section 6.

Here is the article's plan: the data structure of the radiation experiment is described in Section 2. This sets up the main thrust of the article, the efficient reduction of microarray data (320 numbers per gene in this case) to a single summary statistic " Z_i " for each gene, followed by an appropriate simultaneous inference for the activity of each gene based on all the Z scores. Section 3 presents the simple nonparametric empirical Bayes model used to make our simultaneous inferences. The model is presented in algorithmic form, suggesting how it can be applied to other microarray comparative experiments, both oligoneucliotide and cDNA types. (Another such experiment is briefly discussed in Section 6.)

Section 4 concerns the efficient reduction of the data to a single score Z_i per gene. The reduction makes use of the empirical Bayes model, essentially selecting mappings that maximize the amount of Bayesian information preserved in Z_i . Frequentist justification of the empirical Bayes approach appears in Section 5, where it is related to Benjamini and Hochberg's (1995) theory of FDR. Section 6 closes with a summary and some detailed remarks, including a comparison of our analysis with a "gold standard" assay of some of the genes.

2. THE DATA

Microarray experiments produce enormous amounts of data, more than two million feature numbers in the relatively small experiment we are discussing here. The statistical task is to efficiently reduce these numbers to simple summaries of the genes' activities. One goal in this article is to provide a method for comparing the statistical efficiency of different data reduction strategies.

Here is a description of the data in the radiation experiment, and the notation we will use to describe it. Expression levels were recorded for 6,810 different genes,

genes:
$$i = 1, 2, ..., n = 6,810.$$
 (2.1)

(There were actually 7,129 genes, 319 of which had some missing data. For convenience, this article considers only the 6,810 genes having complete data. The various analyses were also carried out on all 7,129 genes, with nearly identical results.) Each gene on each plate was represented by 20 oligonucleotide "probes,"

probes:
$$j = 1, 2, \dots, J = 20$$
. (2.2)

Finally there were eight plates, (the individual microarrays) representing the eight experimental conditions of the experiment described in section 1, (U1A, U1B, I1A, I1B, U2A, U2B, I2A, I2B),

plates:
$$k = 1, 2, ..., K = 8$$
. (2.3)

Two features were recorded for each probe of each gene on each plate, a "perfect match number" pm_{ijk} and a "mismatch number" pm_{ijk} , the latter referring to a deliberately distorted version of the oligonucleotide included as a control. Table 1 shows the 20 pairs of numbers for gene i = 2,715 on plate k = 1.

We will investigate three separate stages of data reduction: "probe reduction", the mapping that takes the 20 probe pair

Probe 2 3 4 5 6 7 8 9 10 pm 1,054 3,242 1,470 4,050 1,356 1,476 561 606 1,307 1,057 793 2,333 826 1,912 561 558 942 526 699 1,060 mm Probe 11 12 13 14 15 16 17 18 19 20 pm 974 1,584 802 1,399 1,670 2.514 2,096 6,592 5.662 2,244 mm 1,771 601 8.717 1,484 668

Table 1. The 20 Pairs of Perfect Match and Mismatch Feature Numbers for Gene i = 2,715 on Plate k = 1 (U1A)

numbers into a single expression value " M_{ik} " for gene i on plate k,

probe reduction:
$$\{(pm_{ijk}, mm_{ijk}), j = 1, 2, \dots, 20\} \rightarrow M_{ik},$$

$$(2.4)$$

"gene reduction," the mapping that takes the K = 8 expression values M_{ik} for gene i into a single expression score " Z_i ,"

gene reduction:
$$\{M_{ik}, k = 1, 2, ..., 8\} \rightarrow Z_i$$
, (2.5)

and finally an *inference mapping* that re-expresses Z_i in terms of a statistical inference concerning i activity of gene. The nonparametric empirical Bayes analysis of Section 3 will provide inferences of the form Prob {Event_i| Z_i }, where Event_i is an event of interest such as "gene i's activity was affected by radiation." Section 5 connects these probabilities with the frequentist FDR criteria of Benjamini and Hochberg (1995).

There are of course an unlimited selection of possible data reductions from the original data, 320 numbers per gene in the radiation experiment, to the expression scores Z_i . For reasons explained in Section 4, the empirical Bayes analysis will lead us to prefer the following choices: For the probe reduction let

$$M_{ik} = \text{mean}\{\log(\text{pm}_{ijk}) - .5 \cdot \log(\text{mm}_{ijk}), j = 1, 2, \dots, 20\}.$$
(2.6)

For the gene reduction, first compute the four differences $(D_{i1}, D_{i2}, D_{i3}, D_{i4})$ between the irradiated and unirradiated values within the same wild-type sample and aliquot, for example

$$D_{i1} = M_{i3} - M_{i1}, (2.7)$$

the difference between the IIA and U1A values M_{ik} . Then take

$$Z_i = \overline{D}_i / (a_0 + S_i), \tag{2.8}$$

where \overline{D}_i is the average of the four differences, S_i is their sample standard deviation, and a_0 is the 90th percentile of the 6,810 S values. Specifications (2.6)–(2.8) will be used as a comparison point in all of our numerical examples. They will be compared with other choices in Section 4, including the current one included in the Affymetrix software.

3. EMPIRICAL BAYES INFERENCES

Besides analyzing the radiation data, our goal here is to provide data analytic techniques useful in a variety of microarray situations. With generality in mind we will avoid highly specified models, relying instead on a simple inference model that

is likely to apply to most comparative experiments: that a gene is either affected on unaffected by the treatment of interest, radiation in our case, giving two possible distributions for the expression score "Z," (2.5). Lee et al. (2000) used a normal theory version of this idea, as, less directly, did Li and Wong (2000). Newton et al. (2000) focused on Gamma models. Here we will avoid parametric assumptions. The resulting nonparametric empirical Bayes analysis, which provides a posteriori probabilities of effect for the various genes, is further justified in Sections 4–6.

Let

$$p_1$$
 = probability that a gene is affected,
 $p_0 = 1 - p_1$ = probability unaffected, (3.1)

and

$$f_1(z)$$
 = the density of Z for affected genes $f_0(z)$ = the density of Z for unaffected genes. (3.2)

Then

$$f(z) = p_0 f_0(z) + p_1 f_1(z)$$
(3.3)

is the mixture density of the two populations. In our situation, we can estimate f(z) directly from the 6,810 expression scores Z_i obtained from the data reduction (2.4) and (2.5).

In the absence of strong parametric assumptions such as normality, model (3.3) is useless without an estimate of the "null density" $f_0(z)$. Fortunately, it is easy to obtain such estimates. What follows is the method we used to estimate $f_0(z)$ in the radiation experiment. Section 6 discusses variants of this method that are applicable more generally.

The 6.810×8 matrix **M** of expression values (2.4), one value for each gene on each plate, gives a 6.810×4 matrix **D** of differences between the irradiated and unirradiated expression values, as in (2.7). Let \mathbf{M}_k indicate the kth column of \mathbf{M} , a 6.810 vector. With the plates ordered as before, (U1A, U1B, I1A, I1B, U2A, U2B, $\mathbf{\Sigma}$ A, $\mathbf{\Sigma}$ B), the "difference matrix" **D** is

$$\mathbf{D} = (\mathbf{M}_3 - \mathbf{M}_1, \mathbf{M}_4 - \mathbf{M}_2, \mathbf{M}_7 - \mathbf{M}_5, \mathbf{M}_8 - \mathbf{M}_6). \tag{3.4}$$

Symbolically, the vector \mathbf{Z} of expression scores (2.5) is obtained via

[original data]
$$\rightarrow$$
 M \rightarrow **D** \rightarrow **Z**. (3.5)
6,810×20×2×8 6,810×8 6,810×4 6,810

Now let the "null difference matrix" **d** be the $6,810 \times 4$ matrix obtained by differencing within the aliquot splits,

$$\mathbf{d} = (\mathbf{M}_2 - \mathbf{M}_1, \mathbf{M}_4 - \mathbf{M}_3, \mathbf{M}_6 - \mathbf{M}_5, \mathbf{M}_8 - \mathbf{M}_7), \tag{3.6}$$

so for example the first column of **d** records differences between the B and A splits of the unirradiated wild-type 1 experiments. We define "null scores" $\mathbf{z} = (z_1, z_2, \dots, z_{6.810})'$ by

[original data]
$$\rightarrow$$
 M \rightarrow **d** \rightarrow **z**, (3.7)

with the understanding that except for the substitution of \mathbf{d} for \mathbf{D} , the arrows in (3.7) indicate the same mappings as in (3.5).

We will use the empirical distribution of the null scores $\{z_i\}$ to estimate the null density $f_0(z)$ in (3.3). One could just as well take $\mathbf{M}_1 - \mathbf{M}_2$ as $\mathbf{M}_2 - \mathbf{M}_1$ in (3.6), etc., and in fact our numerical algorithm employs random sign permutations of the columns of \mathbf{d} to improve the estimation of f_0 . The basic idea here, that we can recover the "null hypothesis" from differences that negate treatment effects, shows up in one form or another in many of the microarray references, being essentially unavoidable in a comparative experiment. Further discussion appears in Section 6, which describes strategies that might be used for estimating f_0 in situations less intricate than the radiation experiment.

An application of Bayes' rule to the mixture model (3.3) gives the a posteriori probabilities $p_1(Z)$ and $p_0(Z)$ that a gene with score Z was affected or unaffected by the treatment:

Bayes' Rule:
$$p_1(Z) = 1 - p_0 f_0(Z) / f(Z)$$

and

$$p_0(Z) = p_0 f_0(Z) / f(Z).$$
 (3.8)

The ratio $f_0(Z)/f(Z)$ can be estimated directly from the $\{Z_i\}$ and $\{z_i\}$ empirical distributions. The probabilities p_0 and $p_1 = 1 - p_0$ are unidentifiable without strong parametric assumptions, but this will turn out to be less problematic than it might seem. The constraint that $p_1(Z)$ be nonnegative for

all Z does restrict p_0 and p_1 ,

$$p_1 \ge 1 - \min_{Z} \{ f(Z) / f_0(Z) \}$$

and

$$p_0 \le \min_{Z} \{ f(Z) / f_0(Z) \} \tag{3.9}$$

A more stable bound for p_1 and p_o is given in Remark F of Section 6.

Figure 1 displays the Bayesian inference curve $p_1(Z)$ = Prob{Event|Z} obtained from the probe and gene data reductions (2.6), (2.8). It was constructed as follows (skipping some technical details that appear in Section 6):

Algorithm 1: Empirical Bayes analysis for microarrays

- (a) Compute the scores $\{Z_i\}$ according to (3.5), using probe reduction (2.6) and gene reduction (2.8).
- (b) Compute the null scores $\{z_i\}$ in the same way, beginning with (3.7). Generate 20 versions of the $\{z_i\}$, based on 20 independent row-wise sign permutations of **d** (see Remark D).
- (c) Use logistic regression to estimate the ratio $f_0(z)/f(z)$ based on the relative densities of the $\{Z_i\}$ to the $\{z_i\}$ (see Remark C).
- (d) Use relationship (3.9) to obtain an estimated upper bound for p_0 : here $p_0 = .811$ (See Remark F.)
- (e) For each gene compute $\operatorname{Prob}\{\operatorname{Event}|Z\}$ from (3.8), with f_0/f estimated from the logistic regression, and p_0 equaling its estimated maximum value (or more conservatively with $p_0=1$.)

We have focussed on our particular experimental setup, but Algorithm 1 is quite general. It can be applied to any twoclass situation, for example two sets of unpaired samples. All that changes is the generation of null scores z_i in step (b). For instance, for unpaired samples the values of z_i would be generated by random permutations of the column labels "1" and "2." Remark E of Section 6 gives another example.

Our Bayesian analysis is actually "empirical Bayes" in the sense that the crucial ratio $f_0(Z)/f(Z)$ in (3.8) is estimated

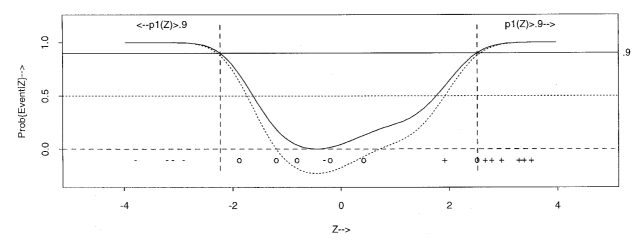


Figure 1. Solid curve: Bayesian Inference Mapping $Prob\{Event_i|Z_i\}$ From Data Reductions (2.6), (2.8); Event_i is "Gene i Affected by Radiation." Symbols show Z values for 18 genes separately analyzed by Northern Blot: "+" positively affected; "-" negatively affected"; "o" not affected. Dotted curve is lower bound (3.10).

from the data rather than from *a priori* assumptions. Newton et al. (2000) carried out a similar analysis, but using specific Bayesian modeling assumptions beyond (3.1)–(3.3).

In Figure 1, the *a posteriori* probability of being affected is seen to increase as Z or -Z grows large. The positive end of the Z-axis corresponds to genes turned *on* by the radiation, with their expression values increased, while negative values of Z indicate decreased expression under radiation. Out of the 6,810 genes 127 genes had $p_1(Z)$ exceeding .90, more on the negative than positive end of the Z scales.

Eighteen of the 6,810 genes were independently assessed by a Northern Blot analysis, a pre microarray assay that serves here as a gold standard for gene expression. Seven of these, indicated by "+" in Figure 1, were deemed "affected positively by radiation," five indicated by "-" were "affected negatively," and six indicated by "o" were "not affected." There is good agreement between the Northern Blot assessments and the probabilities assigned in Figure 1. The full results, given in Section 6, show a high correlation between the gold standard and our results.

In comparing different data reductions, it is convenient to always have the same marginal distribution for Z. To this end, the raw scores $\{Z_i\}$ from (2.8) were monotonically transformed to have a nearly perfect N(0,1) distribution, say by transformation m(Z), and then the null scores were transformed according to the same m(z). Notice that the crucial ratio $f_0(z)/f(z)$ remains the same under such transformations, so that $p_1(Z)$ and $p_o(Z)$ in (3.8) are transformation invariant. We will always make the empirical distribution of the $\{Z_i\}$ almost perfectly N(0,1), using a normal scores transformation, implying for example that $42 = 6,810 \cdot (1-\Phi(2.5))$ of the 6,810 genes have $Z_i > 2.5$, with Φ the standard normal cumulative distribution function.

Figure 2 shows the estimates of f_0 , f_1 , and f contributing to Figure 1; f(Z) is a standard N(0,1) density, by construction, while $f_0(z)$ is a less dispersed density. This is what we hoped for of course: the values of Z should be more dispersed than the values of z because they reflect the disturbing effects of the radiation treatment. The large values of $Prob\{Event|Z\}$ in the tails of Figure 1 come from (3.8), and the small ratio of $f_0(z)$ to f(Z). A good choice of data reductions makes $f_0(z)/f(z)$ small for |z| large, and we will use this criterion to guide our choices of the probe and gene reductions in Section 4.

Looking again at (3.8),

$$p_1(Z) \ge 1 - f_0(Z)/f(Z),$$
 (3.10)

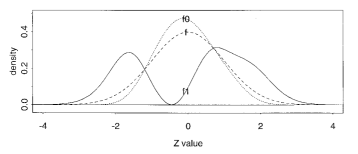


Figure 2. Estimates of $f(\cdot)$, $f_0(\cdot)$ and $f_1(\cdot)$ for the Situation of Figure 1, Model (3.1)–(3.3); $p_0=.811$, its Estimated Maximum From (3.9), Used in the Construction of f_1 .

because this corresponds to $p_0 = 1$, the largest possible value. The dotted curve in Figure 1 is $1 - f_0(Z)/f(Z)$. This is not much less than the solid curve for large values of |Z|, giving 106 genes with $p_1(Z) \ge .90$.

EFFICIENT DATA REDUCTIONS FOR MICROARRAY EXPERIMENTS

The empirical Bayes analysis of Section 3 depends on a drastic data reduction: from the full vector \mathbf{v}_i of data for gene i, a 320-vector in the radiation experiment, to a single number Z_i (and its null counterpart z_i .) Information is bound to be lost in the mapping from \mathbf{v}_i to Z_i , but the less we lose the more powerful will be the analysis, and better our chance of detecting genuinely affected genes.

To state things more exactly, we can imagine applying model (3.1)–(3.2) to the 320-dimensional densities of \mathbf{v} ,

$$f_1^{\mathbf{v}}(\mathbf{v}) = \text{density of } \mathbf{v} \text{ for affected genes,}$$
 (4.1)

 $f_0^{\mathbf{v}}(\mathbf{v}) = \text{density of } \mathbf{v} \text{ for unaffected genes,}$

and

$$f^{\mathbf{v}}(\mathbf{v}) = p_0 f_0^{\mathbf{v}}(\mathbf{v}) + p_1 f_1^{\mathbf{v}}(\mathbf{v}), \tag{4.2}$$

the mixture density; p_0 and p_1 have the same meaning here as in (3.1). Defining the likelihood ratio statistic $R^{\mathbf{v}}(\mathbf{v}) = f^{\mathbf{v}}(\mathbf{v})/f_0^{\mathbf{v}}(\mathbf{v})$, Bayes theorem gives

$$p_1^{\mathbf{v}}(\mathbf{v}_i) = 1 - p_0/R^{\mathbf{v}}(\mathbf{v}_i) = \text{Prob}\{\text{gene } i \text{ affected}|\mathbf{v}_i\}, \quad (4.3)$$

compared to $p_1(Z_i) = 1 - p_0/R(Z_i)$ in (3.8), where $R(Z_i) = f(Z_i)/f_0(Z_i)$.

In our situation it is not practical to estimate the highdimensional densities $f^{\mathbf{v}}(\mathbf{v})$ and $f_0^{\mathbf{v}}(\mathbf{v})$, at least not without extensive modeling. However, we can easily estimate the corresponding densities f(Z) and $f_0(Z)$ for a one-dimensional statistic Z. The goal is to choose a mapping $Z = s(\mathbf{v})$ that does not lose much information. Information loss manifests itself by reductions in the likelihood ratio $R(Z_i)$, compared to $R^{\mathbf{v}}(\mathbf{v}_i)$, which reduces the number of genes having convincingly large values at $p_1(Z_i)$.

4.1 Estimation of a_0

With this background in mind we searched for mappings $Z = s(\mathbf{v})$ that produced large values of R(Z), i.e., good separation between f(Z) and $f_0(Z)$ as in Figure 2. Figure 3 shows the part of the search relating to the choice of a_0 in the denominator of (2.8). The curve marked "90" is equivalent to the dashed curve in Figure 1, the difference here being that the vertical axis is plotted on the logit scale, $\log p_1(Z)/(1-p_1(Z))$, to emphasize differences in the tails. Keeping probe reduction (2.6) fixed, Figure 3 compares five different choices of a_0 in the gene reduction (2.8): a_0 equal to the 90th percentile of the 6,810 S_i values; the 50th percentile; the 5th percentile; $a_0 = 0$; and $a_0 \to \infty$. The choice $a_0 = 0$ makes Z_i in (2.8) proportional to the one-sample t-statistic for the four differences $(D_{i1}, D_{i2}, D_{i3}, D_{i4})$, whereas $a_0 \to \infty$ makes Z_i equivalent to the numerator D_i . The plotted curves

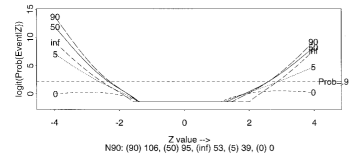


Figure 3. Choice of a_0 in the Gene Mapping $Z_i = \overline{D_i}/(a_0 + S_i)$, (2.8); Vertical Axis is Logit of Prob{Event|Z}, Estimated as in (3.8) With $p_0 = 1$; "90" Indicates a_0 Equaling 90th Percentile of the 6,810 S_i Values, etc.; "inf" is Limit as $a_0 \to \infty$. We see that 90 is the best choice in terms of maximizing Prob{Event|Z} for large |Z|; $a_0 = 0$ is worst. All choices used probe reduction (2.6). The vertical axis is truncated at lower bound Prob{Event|Z} = .20. **N90** is the number genes having Prob{Event|Z} \geq .90.

are the logits of (3.10), the conservative lower bound for $p_1(Z)$, (taking $p_0 = 1$ in (3.8).)

Figure 3 shows that the best choice for a_0 is the one we used before, a_0 the 90th percentile. This manifests itself as higher values of Prob{Event|Z} at both ends of the Z scale. The density $f_0(z)$ in Figure 2 is more concentrated around zero than it is say for the disastrous choice $a_0 = 0$, raising $f(Z)/f_0(Z)$ in the tails and thus $p_1(Z)$, (3.10). The numbers N90 in Figure 3 indicate the number of genes having lower bound (3.10) for $p_1(Z_i)$ greater than .90. These range downward from 106 for $a_0 = 90$ to 0 for $a_0 = 0$. Larger values of N90 indicate less information loss in going from the full data vector \mathbf{v}_i to the summary statistic Z_i . (Efron, Tibshirani, Goss, and Chu (2001) also use Kulback–Liebler distance to measure information loss.)

1: c=.5 & logs N90: 106 2: c=1 no logs N90: 36 3: c=.5 no logs N90: 62 4: c=0 no logs N90: 14

4.2 Choosing the Probe Reduction

The GeneChip software distributed by Affymetrix uses a simple average difference, (with some outlier rejection) to estimate what we called the probe reduction in (2.4), the expression for gene i on plate $k: M_{ik} = \text{mean}_j \{ \text{pm}_{ijk} - \text{mm}_{ijk} \}$. However, this choice is controversial, and some researchers have suggested that ignoring the mismatch entirely might produce better expression estimates. We investigate the issue here.

Keeping the gene reduction fixed as in (2.6), $a_0 = .90$, Figure 4 compares probe reductions of the form

$$M_{ik} = \operatorname{mean}_{i} \{ s(\operatorname{pm}_{iik}) - c \cdot s(\operatorname{mm}_{iik}) \}, \tag{4.4}$$

with s either the log function or the identity function. For example curve 2 in the left panel uses $M_{ik} = \text{mean}_j \{\text{pm}_{ijk} - \text{mm}_{ijk}\}$ whereas the dotted curve in the right panel uses $M_{ik} = \text{mean}_j \{\log(\text{pm}_{ijk})\}$. Our preferred choice (2.6)–(2.8) is curve 1, "c = .5 & logs." The "Affy" curve in the left panel was based on the algorithm provided by Affymetrix, which is similar to the "c = 1 no logs" choice, but with a provision for removing apparent outliers among the 20 pm_{ijk} – mm_{ijk} differences before averaging.

Figure 4 indicates a substantial advantage to taking logs, and a mild advantage to using c=.5 rather than c=1 or c=0. The comparison between c=.5 and c=1 is close on the log scale, but other comparisons, reported in Efron et al. (2000), reinforce the superiority of c=.5. We also tried using various L-estimators in (4.7), including trimmed means. When applied on the log scale, this form of robustification made almost no difference to our results.

Some comments are in order, which apply to the whole section:

• There is no claim that the mapping $Z = s(\mathbf{v})$ described by (2.6), (2.8) is "correct," only that it is relatively efficient in preserving the information in \mathbf{v} . The estimated curve

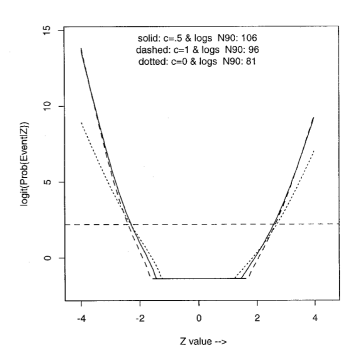


Figure 4. Comparison of Various Probe Reductions (gene reduction fixed as in (2.8), $a_0 = 90$ th percentile). The solid curve in both panels is the choice (2.6) used previously; constant "c" is multiple of mm level subtracted from pm level, for example. "c = 1 no logs" uses $M_{ik} = mean\{pm_{ijk} - mm_{ijk}\}$. "Affy" based on the probe reduction software provided with the Affymetrix Genechip.

- $p_1(Z)$ still is meaningful as the *a posteriori* probability of effect given the insufficient statistic Z, and also has the FDR interpretation of Section 5. (Efron et al. (2000) showed that in fact a better " $s(\mathbf{v})$ " can be obtained in the radiation experiment by removing plates U1A and I1A from the dataset (1.1); a processing error appears to have degraded the results from I1A.) Our tactic of choosing the Z mapping to maximize $p_1(Z)$ is nearly equivalent to minimizing FDR, which was the approach taken in Tusher et al. (2000).
- There is also no claim that mappings (2.6) and (2.8) enjoy general superiority. The equivalent of Figures 3 and 4 might point to a different choice of $s(\mathbf{v})$ in another dataset. Section 6 discusses how our methodology can be applied to other comparative microarray experiments.
- Overfitting is not a threat in a genuine Bayesian framework, where results like those from Figures 3 and 4 can be thought of as just computer-based attempts to numerically solve a probabilistic maximization problem. However, in our empirical Bayes framework, too much data-based maximization could in fact lead to overfitting. Two forms of bootstrapping were employed as a check on our results: "gene resampling," in which the rows of the 6,810 × 8 matrix M were resampled to give M*; and "row resampling" in which row i of M* was obtained as the average of 20 resampled rows from the 20 × 8 matrix x_i having entries

$$x_{ijk} = \log(\text{pm}_{ijk}) - .5 \cdot \log(\text{mm}_{ijk}). \tag{4.5}$$

The bootstrap results indicated that the differences seen in Figures 3 and 4 were much greater than the standard errors of the curves, so that overfitting was not a threat. For example, row resampling showed that the difference between the $a_0=.90$ and $a_0=.50$ curves at Z=-3, which looks suspiciously small in Figure 3, had point estimate and standard error $.68\pm.13$.

5. FALSE DISCOVERY RATES

The empirical Bayes analysis of Section 3 is closely related to Benjamini and Hochberg's FDR criterion. For a collection of simultaneous hypothesis tests, FDR is the expected proportion of type I errors made using a given rejection rule. Define the *local false discovery rate* to be

$$fdr(Z) = p_0 f_0(Z) / f(Z),$$
 (5.1)

so fdr(Z) is the *a posteriori* probability $p_0(Z)$, (3.8), that a gene with score Z is unaffected. It will be shown that (6.1) has a natural FDR interpretation. We begin with a numerical example.

In the calculations for Figure 1, N = 74 of the 6810 genes had Z scores in the interval $Z \in [1.9, 2.1]$, whereas the twenty permuted null score datasets $\{z_i\}$ had 676 falling into [1.9, 2.1], an average of 33.8 = 676/20 per set. Taking p_0 to be its estimated maximum .811, this suggests that among the N = 74 binned Z values, the expected number of "unaffecteds" is $27.4 = .811 \cdot 33.8$. If we now declare all genes with Z in [1.9, 2.1] to be affected, our expected proportion of false discoveries is

$$\frac{27.4}{74} = 37\%. \tag{5.2}$$

Notice that (5.2) is an obvious estimator of (5.1) for Z = 2,

$$\widehat{\text{fdr}}(2) = \hat{p_0} \hat{f_0}(2) / \hat{f}(2)$$

$$\left[\hat{p_0} = .811, \quad \hat{f_0}(2) = \frac{33.8}{6810}, \quad \hat{f}(2) = \frac{74}{6810} \right]. \quad (5.3)$$

In general, if we bin the genes into small intervals on the Z scale, then a bin declared "affected" will have a FDR of about fdr(Z), (5.1), the equality becoming exact as the number of genes goes to infinity. This last statement can be rigorously verified under modest ergodic conditions that preclude extremely high correlations among the values fo Z or the z.

Figure 5 reports on a simulation experiment used to check the accuracy of fdr(Z) as an estimate of FDR. A 6810×8

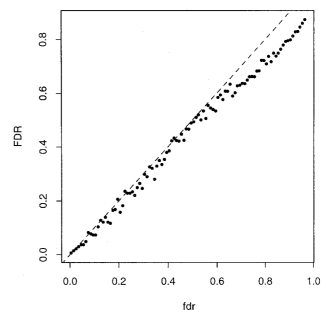


Figure 5. Simulations Comparing Empirical Bayes Formula fdr, (5.1) With Actual FDR, as Explained in Text. Left panel: $a_0 = .90$ in mapping (2.8). Right panel: $a_0 \to \infty$.

matrix M was constructed in a way that mimicked the radiation experiment,

$$M_{ik} = \theta_i t_k + \epsilon_{ik} \quad [\epsilon_{ik} \stackrel{\text{ind}}{\sim} N(0, 2)],$$
 (5.4)

 $(t_1, t_2, \ldots, t_8) = (0, 0, 1, 1, 0, 0, 1, 1);$ 681 of the "gene effects" θ_i were chosen from an N(-1.5, 1) distribution, 681 from N(1.5, 1), and the remaining 5448 set at zero. In other words 80% of the genes were unaffected and 20% were affected, 10% in each direction.

The matrix **M** was processed into a **Z** vector according to (3.5), and also into twenty **z** vectors according to (3.7), using (2.8) for the mappings from $\mathbf{D} \to \mathbf{Z}$ and $\mathbf{d} \to \mathbf{z}$. Following the same algorithm that lead to Figure 1, these gave an estimated $\mathrm{fdr}(Z)$ curve, (5.1), that in fact looked much like the one for the actual experiment.

Figure 5 reports on two different choices of a_0 in (2.8), $a_0 = .90$ in the left panel and $a_0 \rightarrow \infty$ on the right. There are 100 points in each panel, corresponding to a binning of the fdr axis in units of .01 from 0 to 1, with the *j*th point plotted at fdr_i = j/100 and

$$FDR_i = \text{proportion of } j\text{th bin with } \theta_i = 0,$$
 (5.5)

the empirical FDR for that bin. The FDR_j values are averages over 50 simulations, each of the individual simulations being noisy versions of the same picture. If formula (5.1) is actually the local fdr, then the points should lie near the main diagonal FDR = fdr as they do. The slight conservative bias FDR_j \leq fdr_j, came from the fact that the upper bound for p_0 used in (5.1) (calculated as in Remark F of Section 6) substantially overestimated the true value $p_0 = .80$.

In the artificial situation (5.4), taking $a_0 \rightarrow \infty$ in (2.8) gives a more efficient choice of Z_i than $a_0 = .90$, doubling N90. Nevertheless, the inefficient choice $a_0 = .90$ is still accurately calibrated: $FDR_i \doteq fdr_i$.

FDRs are usually defined for an entire rejection region, for example for

$$\mathcal{R} = \{Z: R(Z) \ge r_0\} \quad [R(Z) = f(Z)/f_0(Z)] \tag{5.6}$$

rather than locally as in (5.1). We can think of this as replacing our original choice of summary statistic $Z = s(\mathbf{v})$ with

$$\widetilde{Z} = \begin{cases} 1 & Z \in \mathcal{R}, \\ \widetilde{Z} = & \text{if} \\ 0 & Z \notin \mathcal{R} \end{cases}$$
 (5.7)

Assuming that genes having $\widetilde{Z} = 1$ are declared affected, the empirical Bayes formula (5.1) now becomes

$$\widetilde{\text{fdr}}(1) = p_0 \tilde{f}_0(1) / \tilde{f}(1),$$
 (5.8)

with straightforward estimate

$$\hat{p_0} \frac{\text{proportion}\{z_i \in \mathcal{R}\}}{\text{proportion}\{Z_i \in \mathcal{R}\}}.$$
 (5.9)

The heuristic argument proceeding (5.2) is more obvious here: (5.9) estimates the proportion of genes in the "affected" region \mathcal{R} , which is actually unaffected, and in this sense estimates

Benjamini and Hochberg's FDR. (Current work by the authors strengthens this connection: choosing \mathcal{R} in (5.6) as large as possible subject to keeping (5.9) below some fixed limit exactly matches the Benjamini–Hochberg choice of rejection region.)

The global definition of FDR has the advantage of being easier to estimate. We can use the totally nonparametric estimator (5.9) rather than having to estimate the local ratio $f_0(Z)/f(Z)$ in (5.1). On the other hand (5.8) is a composite measure that assigns the same FDR to all the genes in $\mathcal R$ even though some of them have $\operatorname{Prob}\{\operatorname{Event}|Z\}$ much greater than others. Comparing (5.1) with (5.8) it is easy to see that $\operatorname{fdr}(1)$ is the conditional expectation of $\operatorname{fdr}(Z)$ given $Z \in \mathcal R$,

$$\widetilde{\mathrm{fdr}}(1) = E_f\{\mathrm{fdr}(Z)|\mathcal{R}\}\tag{5.10}$$

so that fdr(Z) is more precise than $\widetilde{fdr}(1)$. More on the connection between FDR and fdr appears in Storey (2001).

6. SUMMARY AND REMARKS

The Empirical Bayes procedure described in this article provides an effective framework for studying the relative changes in gene expression for a large number of genes. It uses a simple nonparametric mixture prior to model the population of affected and unaffected genes, thereby avoiding parametric assumptions about gene expression. We establish a close connection between the estimated posterior probabilities and a local version of the FDR, thereby allowing for the analyst to handle multiple testing issues that arise when dealing with a large number of simultaneous tests. As we have detailed in Algorithm 1 and Remark D, the proposed procedure can be applied quiet generally to other kinds of comparative microarray experiments.

We conclude with a number of remarks, giving important practical details for the proposed methods.

(A) The Experiment. Lymphoblastoid cell lines GM14660 and GM08925, (Coriell Cell Repositories, Camden, New Jersey) were seeded at 2.5×105 cells/ml. The treatment consisted of 5 Gy of ionizing radiation. After 24 hours, RNA was isolated, labeled, and divided into two aliquots that were independently hybridized to the HuGeneFL Genechip microarray, Affymetrix Corporation.

(B) Northern Blot Analysis. Northern Blot Analysis produced a quantitative score " G_i " for each of the 18 genes indicated in Figure 1, G standing for gold standard. Following previous biomedical convention, G scores exceeding 1.30 were taken to indicate a positive effect of radiation on gene activity, the "+" symbols in Figure 1; likewise "-" for $G_i < .70$ and "o" for $.70 \le G_i \le 1.30$. Figure 6 compares the Z_i scores from Figure 1 with $\log G_i$ for the 18 test genes. We see a strong monotone relationship, correlation .87.

The agreement in Figure 6 is impressive, especially considering the magnitude of the sampling errors in the individual expression values. Our gold standard, the Northern Blot score, is not pure gold, itself being subject to experimental error. There is only one flagrant disagreement in Figure 6, the "-" gene at $Z_i = -.31$. The vector of differences (3.4) was $\mathbf{D}_i = (-1.59, .55, ,88, -.83)$ for this gene, so that both wild types yielded aliquots of opposing signs. In contrast the "o"

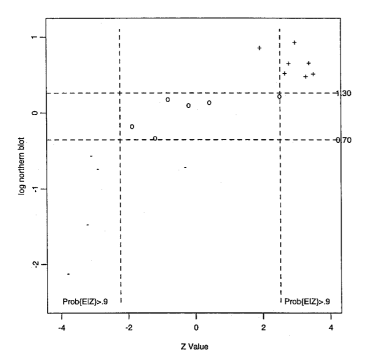


Figure 6. Comparison of Z Scores From the Analysis in Figure 1 With the Logarithm of the Northern Blot Results. Correlation .87. Z values outside the two vertical lines have $Prob\{Event_i \mid Z_i\} \ge .90$.

point at $Z_i = 2.51$, lying just below the "+" cutoff value G = 1.30, was consistently positive, $\mathbf{D}_i = (4.54, 2.81, 1.64, 2.65)$, strengthening our belief that this gene was positively affected by the radiation.

(C) Debrightening and Desumming. Some microarray plates are "brighter" than others in that they produce systematically larger expression levels. Following probe reduction (2.4) we debrightened the data by separately standardizing the columns of M. That is, each column of M was linearly transformed to have mean 0 and empirical standard deviation 1.

"Desumming" corrects for another type of data inhomogeneity. Corresponding to \mathbf{D} (3.4), let

$$S = (M_3 + M_1, M_4 + M_2, M_7 + M_5, M_8 + M_6).$$
 (6.1)

A gene with larger **S** values tended to have larger values of **D**, which undercut the exchangeability across genes implicit in our empirical Bayes analyses. (Newton et al. 2001 adjusted their data for a similar problem.) After debrightening, the individual columns of **D** were desummed as follows: a linear regression $|D_{ik}| = a_0 + a_1 |S_{ik}| + \text{error}$ was fit individually to each column, and then each D_{ik} was transformed to

$$D_{ik}/(\hat{a_0} + \hat{a_1}|S_{ik}|).$$
 (6.2)

Similar transformations were made on the columns of \mathbf{d} , (3.6). It was the transformed \mathbf{D} and \mathbf{d} matrices that were used to compute the scores \mathbf{Z} and \mathbf{z} via (2.8). Desumming made almost no difference to the results in Figure 1, but the exchangeability issue is an important general point of concern for the empirical Bayes analysis, see Remark F.

(D) Logistic Regression Estimate of $f_0(z)/f(z)$. The ratio $f_0(z)/f(z)$ in (3.8) was estimated by logistic regression. Given B=20 replications of \mathbf{z} , all $n \cdot (1+B)=6$, $810 \cdot 21$ scores Z_i

and z_i were plotted on a line, with values of Z_i considered as "successes" and values of \mathbf{z} , as "failures". The probability $\pi(z)$ of a success at point z is given in terms of the densities (3.2),

$$\pi(z) = f(z)/(f(z) + Bf_0(z)), \tag{6.3}$$

so that (3.8) becomes

$$p_1(Z) = 1 - p_0 \frac{1 - \pi(Z)}{B\pi(Z)}. (6.4)$$

With n = 6,810 genes, the normal scores transformation resulted in $\max\{Z_i\} = -\min\{Z_i\} = 3.80$, whereas the null scores $\{z_i\}$ were confined to a smaller range, as in Figure 2. Our algorithm divided the range [-4, 4] into 139 equal intervals, counted the number of values of Z_i and z_i in each interval, and estimated $\pi(z)$ by logistic regression, for use in (6.4). The regression function was a natural spline with 5 degrees of freedom, called by the Splus command ns(x, df = 5), x being the 139 center points of the intervals. The choice of B = 20 z replications was based on an analysis like that in Figure 3, which showed considerable improvement for B increasing from 1 to 10, but little gain past 20. Other methods of estimating $f_0(z)/f(z)$ are possible, and in fact the details of the logistic regression method made little difference to our results. The "global" estimate (5.9) avoids regression entirely, at the expense of providing less-specific results.

(E) Estimating the Null Distribution. The null density $f_0(z)$ is supposed to describe the distribution of expression scores for genes unaffected by the treatment of interest. Basing f_0 on **d** in (3.6) seems natural for the radiation experiment, but other choices are possible and may be necessary for other experimental designs. Table 2 shows a small portion (5 of 2,638 genes) of the data from a microarray study comparing two different types of liver cancer, 36 Type I patients versus 36 Type II, with Type II having worse prognosis. Spotted cDNA arrays were used, the "red-green" variety, the tabled values being $10^3 \cdot \log$ (red/green) intensity ratio. The full table corresponds to matrix M in (3.5), now $2,638 \times 72$. Probe reduction (2.8) is very simple here, (red, green) $\rightarrow \log(\text{red/green})$, though more efficient reductions may be possible as shown in Dudoit et al. (2000). The analog of Figure 3 indicated a preference for $a_0 = 0$, i.e. for taking Z to be the two-sample t-statistic between the Types.

We obtained the null scores z_i and the null density $f_0(z)$ by randomly splitting the Type I patients into two groups of 18 each, say "A" and "B," and likewise "C" and "D" for the Type II patients, and defining the values of z as t-statistics between groups $A \cup C$ versus $B \cup D$. In other words, we used balanced permutations that put equal numbers of those of Type I and Type II into each of the two permuted groups; using unbalanced permutations would add an unwanted component of variance to the null scores. The empirical Bayes analysis produced results similar to those in Figure 2, with Type II playing the role of the Treatment group.

As a simple, but informative, model for the radiation experiment, suppose that M_{ik} in (3.5) can be expressed as

$$M_{ik} = \mu_i + \alpha_i w_k + \theta_i t_k + \epsilon_{ik}, \tag{6.5}$$

	TYPE I						TYPE II				
	pat1	pat2	pat3	pat4	pat5		pat37	pat38	pat39	pat40	pat41
GENE1	230.0	-1,350	−1,580 . 0	-400	-760		970	110	-50	-190.0	-200
GENE2	470.0	-850	8	-280	120		390	-1,730	-1,360	8	-330
GENE3	-920.0	-1,070	1,360.0	-510	-1,120		70	-1,150	340	-400.0	580
GENE4	.1	380	730.0	180	-90		1,040	180	1,070	250.0	1880
GENE5	390.0	-1,960	-210.0	200	230	• • •	530	-1,170	670	.7	890

Table 2. Some Data From a Microarray Study Comparing two Types of Liver Cancer

where $\mathbf{t}' = (0, 0, 1, 1, 0, 0, 1, 1)$, $\mathbf{w}' = (-1, -1, -1, -1, 1, 1, 1, 1)$, and ϵ_{ik} is an independent noise term. Here θ_i represents the treatment effect whereas α_i is the differential response for gene i between the first and second wild types. Then Z_i in (2.8) is

$$Z_i = (\theta_i + e_i)/(a_0 + S_i), \quad S_i = \left[\sum_{\ell=1}^4 (e_{i\ell} - e_i)^{2/3}\right]^{1/2}, \quad (6.6)$$

where each $e_{i\ell}$ is the difference of two values of ϵ_{ik} and e_i is the average of the four values of $e_{i\ell}$. The values of z_i have the same expression except that $\theta_i = 0$ in (6.6). We can see that $f_0(z)$ is a legitimate null hypothesis comparator for f(Z).

Suppose we had defined null scores by differencing across wild types instead of across aliquots: $\mathbf{d} = (\mathbf{M}_5 - \mathbf{M}_1, \mathbf{M}_6 - \mathbf{M}_2, \mathbf{M}_7 - \mathbf{M}_3, \mathbf{M}_8 - \mathbf{M}_4)$ replacing (3.6). Then z_i would pick up an additional term due to the gene/wild-type interaction α_i in (6.5), adding a component of variance to $f_0(z)$, and decreasing the likelihood ratio $f(z)/f_0(z)$. Models like (6.5) are helpful in guiding the choice of the Z and z mappings, even if we do not need them for the data-based estimation of f and f_0 .

The additive model (6.5) gives every column of **d** the same distribution, but we might not trust the Treatment differences to really have the same distribution as the Control, $\square B - \square A$ compared to $\square U2B - \square U2A$ for example. Empirically this turned out not to be a problem for the radiation experiment, but if it had we might have used only the first and third columns of **d** in (3.6).

(F) Better Upper Bound Estimates for " p_0 ." The upper bound (3.9), $p_0 \le \min\{f(Z)/f_0(Z)\}$, can be poorly estimated by the choice $\min\{\hat{f}(Z)/\hat{f}_0(Z)\}$ used in Figure 1. More stable upper bounds can be constructed by integrating over an interval " \mathcal{A} " near Z=0,

$$p_0 \le \frac{\int_{\mathcal{A}} [f(Z)/f_0(Z)] f_0(Z)}{\int_{\mathcal{A}} f_0(Z)} = \frac{\int_{\mathcal{A}} f(Z)}{\int_{\mathcal{A}} f_0(Z)}.$$
 (6.7)

Simulation showed that the choice $\mathcal{A} = [-.5, .5]$ performed better than $\min\{f(Z)/f_0(Z)\}$, particularly when the true p_0 was near 1. The upper bound (6.7) is directly estimated by proportion{values of Z_i in \mathcal{A} }/proportion{values of z_i in \mathcal{A} },

avoiding the logistic regression estimate for $f(Z)/f_0(Z)$. This gave $p_0 \le .825$ in the context of Figure 1, not much different than the previous estimate $p_0 \le .811$. [Received 8 November,

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