Controlling False Discoveries in Multidimensional Directional Decisions, with Applications to Gene Expression Data on Ordered Categories

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SUMMARY: Microarray gene expression studies over ordered categories are routinely conducted to gain insights into biological functions of genes and the underlying biological processes. Some common experiments are time-course/dose-response experiments where a tissue or cell-line is exposed for different doses and/or durations of time to a chemical. A goal of such studies is to identify gene expression patterns/profiles over the ordered categories. This problem can be formulated as a multiple testing problem where for each gene the null hypothesis of no difference between the successive mean gene expressions are tested and further directional decisions are made if it is rejected. Much of the existing multiple testing procedures are devised for controlling the usual false discovery rate (FDR) rather than the mixed directional FDR, the expected proportion of Type I and directional errors among all rejections. Benjamini and Yekutieli (2005) proved that an augmentation of the usual Benjamini-Hochberg (BH) procedure can control the mixed directional FDR while testing simple null hypotheses against two-sided alternatives in terms of one dimensional parameters. In this article, we consider the problem of controlling the mixed directional FDR involving multidimensional parameters. To deal with this problem, we develop a procedure extending that of Benjamini and Yekutieli based on the Bonferroni test for each gene. A proof is given for its mixed directional FDR control when the underlying test statistics are independent across the genes. The results of a simulation study evaluating its performance under independence as well as under dependence of the underlying test statistics across the genes relative to other relevant procedures are reported. Finally, the proposed methodology is applied to a time-course microarray data obtained by Lobenhofer et al. (2002). We identified several important cell-cycle genes, such as DNA replication/repair gene MCM4 and replication factor subunit C2, which were not identified by the previous analyses of the same data by Lobenhofer et al. (2002) and Peddada et al. (2003). Although some of our findings overlap with previous findings, we identify several other genes that compliment the results of Lobenhofer et al. (2002).

KEY WORDS: Benjamini-Hochberg procedure; Directional FDR; Dose-response; Microarray; Multiple testing; Ordered categories; Time-course.

1. Introduction

In many applications researchers are interested in identifying trends in mean response over ordered categories in large scale experiments. With the advent of microarray technology such experiments are common in the literature where investigators are routinely conducting experiments to investigate changes in mean gene expressions over time or dose of a chemical or cancer stage, etc. For example, Lobenhofer et al. (2002) studied the effect of $17-\beta$ estradiol on the gene expression of MCF-7 breast cancer cells as the cells progressed through various phases of cell division cycle. In another experiment, Tamoto et al. (2004) investigated the changes in gene expression with tumor progression in esophagal cancer and identified genes implicated in the early stages of esophagal squamous cell carcinoma. Recently, Bochkina and Richardson (2007) discussed the analysis of a time-course gene expression data where cells from the H2Kb muscle cell line of mouse were treated by insulin (0, 2 or 12 hours of exposure).

In studies such as those described above, identification of statistically significant genes that have similar mean expression profiles over ordered categories is often an important goal to researchers. By identifying such genes, the researchers may potentially discover co-regulated genes belonging to similar pathways and gain insights into biological functions and processes of groups of genes with similar patterns of expressions.

Peddada et al. (2003) introduced an order restricted inference based method for identifying significant genes and group them according to various patterns of inequalities. Implicitly in their methodology, two decisions are being taken for each gene. First, it is decided whether or not a gene is significant using a method exercising a control over gene specific Type I error rate. Then, a suitable inequality pattern is assigned for each selected significant gene based on the values of the underlying test statistics. The directional error that can potentially occur in addition to the usual Type I error due to assigning wrong inequality pattern to a selected

significant gene has not been addressed in that paper. In this paper, we take care of both the Type I and directional errors. We do that by taking a multiple testing approach to the main problem where for each gene the mean expressions are successively compared across the ordered categories and a null hypothesis signifying no particular directional pattern is formed to test against the union of all possible directional patterns. We develop a method of simultaneously testing these null hypotheses and determining a directional pattern upon rejection of each of them controlling both the Type I and directional errors in an overall sense.

Although directional error has not been discussed extensively in the literature, it is perhaps a common error that occurs in applications. While testing a null hypothesis $H_0: \theta = 0$ against the two-sided alternative $H_1: \theta \neq 0$, for some single parameter θ of interest, researchers commonly conclude either $\theta > 0$ or $\theta < 0$ upon rejection of H_0 depending on the sign of the underlying test statistic, keeping the directional error controlled in addition to the Type I error. However, when multiple hypotheses are tested and the number of parameters describing directional patterns is larger, even moderately, than one, as is the case with time course or dose-response microarray data, controlling the directional errors is a problem.

A traditional approach to dealing with directional as well as Type I errors from a multiple testing point of view is to apply a method that controls the so called mixed directional familywise error rate (mdFWER), which is the probability of one or more Type I or directional errors, a variant of the classical familywise error rate (FWER) (Finner, 1994, 1999; Liu, 1996; Sarkar et al., 2004; Shaffer, 1980). However, when the number of null hypotheses is large, as in the context of microarray experiments, the notion of mdFWER, just like the FWER, is too stringent, allowing little chance to make true directional as well as nondirectional discoveries. The FDR (False Discovery Rate), due to Benjamini and Hochberg (1995), is a more powerful concept of overall Type I error rate than the FWER in the context of multiple testing and is now most commonly used in large scale scientific investigations, especially in microarray gene expression studies. A variant of it while controlling both Type I and directional errors would be more powerful than the mdFWER. Two such variants have been introduced in the literature (Benjamini et al., 1993), the pure directional FDR which is the expected proportion of directional errors among rejected hypotheses and the mixed directional FDR (mdFDR) which is the expected proportion of Type I and directional errors among rejected hypotheses. In this article, we will focus on procedures controlling the mdFDR.

Benjamini and Yekutieli (2005) gave a method with independent tests that controls the mdFDR when testing multiple simple hypotheses against two-sided alternatives. They proved that the original Benjamini and Hochberg (1995) procedure controlling the FDR at α can be augmented to make directional decision upon rejecting a null hypothesis according to the value of the corresponding test statistic without causing the mdFDR to exceed α , a result conjectured before by several authors (Benjamini and Hochberg, 2000; Shaffer, 2002; William et al., 1999). Throughout the paper we shall denote Benjamini and Hochberg procedure by BH procedure. Clearly, this method, referred to as the directional BH procedure, can be applied to analyze dose-response microarray data if there are only two ordered categories, but often this is not the case, as such data typically involve more than two ordered categories.

We extend the BH directional FDR procedure in this article to develop our proposed multiple testing method that allows us to make a decision on the directional pattern involving multiple parameters once a null hypothesis of no pattern is rejected and maintains a control over the mdFDR. The proposed methodology is then evaluated through a simulation study and applied to the time-course microarray data in Lobenhofer et al. (2002). Our analysis of Lobenhofer's data resulted in the discovery of several cell-cycle genes that were not previously identified by Lobenhoer et al. (2002) and Peddada et al. (2003). Some of our findings complement the previous findings as detailed in Section 5. An important and unique feature of our methodology is that it permits us to specify the time interval of up (or down) regulation of a gene during the 48 hour period of the cell-cycle. One of the usual objectives for conducting cell-cycle time course experiments is to determine the phase of peak expression for a cell-cycle gene and our methodology allows us to make such determinations.

2. Notations, Definitions and Problem Formulation

In this section, we present the multiple testing formulation of the problem of identifying expression patterns/trends over ordered categories simultaneously for all the genes, having introduced some notations and definitions related to multiple testing.

Let μ_{ij} denote the mean response of the j^{th} variable (e.g. gene), $j = 1, \ldots, m$, in the i^{th} ordered category, $i = 1, \ldots, p$. A problem of biological interest is to group genes by the inequalities among the mean responses, known as directional patterns or order restrictions. Some common inequalities of interest are $\mu_{1j} \leq \mu_{2j} \leq \ldots \leq \mu_{pj}$ (monotone pattern), $\mu_{1j} \leq \mu_{2j} \leq \ldots \leq \mu_{ij} \geq \mu_{(i+1)j} \geq \ldots \geq \mu_{pj}, i = 2, \ldots, p-1$ (umbrella order with peak μ_{ij}). Let $\delta_{ij} = \mu_{i+1j} - \mu_{ij}, i = 1, \ldots, p-1, j = 1, \ldots, m$. Then, the above inequalities of interest or any other inequalities can be restated in terms of the signs of the δ_{ij} 's. Let $\delta_j = (\delta_{1j}, \ldots, \delta_{qj})'$, where q = p - 1. Suppose we test

$$H_{0i}: \boldsymbol{\delta}_{i} = \mathbf{0} \text{ against } H_{1i}: \boldsymbol{\delta}_{i} \neq \mathbf{0}, \tag{1}$$

and decide for a rejected H_{0j} which component δ_{ij} 's are non-zero before declaring their signs to be positive or negative depending on the values of the corresponding test statistics. The declared signs of the δ_{ij} 's then determine a possible inequality or directional pattern. For instance, in the case of q = 4, suppose for a given gene j, the $\delta_j = (\delta_{1j}, \ldots, \delta_{4j})$ is found significantly different from a null vector, with δ_{1j} and δ_{2j} declared to be positive and negative, respectively, and δ_{3j} and δ_{4j} zeros. Then, the corresponding directional pattern is $\mu_{1j} < \mu_{2j} < \mu_{3j} = \mu_{4j} = \mu_{5j}$. We can test H_{0j} against H_{1j} for all the genes applying a suitable multiple testing method. Thus, given p ordered categories for each gene, the task of identifying directional patterns of the mean expressions over these categories for all the genes is being formulated as a multiple testing problem where H_{0j} is tested against H_{1j} simultaneously for all the genes and the signs of the δ_{ij} 's are determined subsequent to the rejection of the corresponding H_{0j} .

For multiple testing of H_{0j} against H_{1j} , j = 1, ..., m, we need *p*-values that will provide a valid test for each of these individual testing problems and will allow us to make decisions on the individual δ_{ij} 's once a H_{0j} is rejected. For that, we consider for each *j* the *p*-value available for testing each component null hypothesis $H_{0j}^i : \delta_{ij} = 0$ against the corresponding component alternative hypothesis $H_{1j}^i : \delta_{ij} \neq 0$, for i = 1, ..., q, and apply a suitable combination method pooling these *q p*-values by treating H_{0j} as an intersection of the subfamily of these *q* component null hypotheses, that is, $H_{0j} = \bigcap_{i=1}^{q} H_{0j}^i$, and H_{1j} as a union of the corresponding *q* alternative hypotheses, that is, $H_{1j} = \bigcup_{i=1}^{q} H_{1j}^i$. Before we discuss appropriate combination methods to be used, let us explain how to obtain these component *p*-values and state the underlying assumptions.

For every $i = 1, \dots, q$ and $j = 1, \dots, m$, suppose we use the absolute value of a test statistic T_{ij} for testing H_{0j}^i against H_{1j}^i . Let $T_{ij} \sim F_{ij}(t, \delta_{ij})$ for some continuous cdf F, which is symmetric about 0 under H_{0j}^i and gets stochastically larger or smaller as δ_{ij} either increases or decreases from 0. In other words, with $F_{ij}(t, \delta_{ij})$ denoting the cdf of T_{ij} at tunder the parameter δ_{ij} , we have $F_{ij}(t, \delta_{ij}) \leq \text{ or } \geq F_{ij}(t, 0)$ according as $\delta_{ij} > \text{ or } < 0$, and $F_{ij}(0,0) = \frac{1}{2}$. Under this setting, a right-tailed test based on the absolute value of T_{ij} will be considered for testing H_{0j}^i against H_{1j}^i , with the corresponding two-sided p-value being defined as $\tilde{P}_{ij} = 2 \min \{F_{ij}(T_{ij}, 0), 1 - F_{ij}(T_{ij}, 0)\}$. By the assumed distributional property of T_{ij} , it is easy to verify that under H^i_{0j} , the two-sided *p*-value \tilde{P}_{ij} satisfies

$$Pr\{\widetilde{P}_{ij} \leq p\} \leq p, \quad \text{for any} \ p \in (0,1).$$
 (2)

Given p-values for testing H_{0j}^i against H_{1j}^i , for $i = 1, \ldots, q$, a number of combination methods (or methods of pooling the p-values) are available in the literature for testing the intersection null hypothesis $H_{0j} = \bigcap_{i=1}^q H_{0j}^i$ against the alternative $H_{1j} = \bigcup_{i=1}^q H_{1j}^i$. Among these, however, the Bonferroni and Simes methods are often used in multiple testing and allow one to make decisions on the individual δ_{ij} 's. For a review of these methods, one may see Bernhard et al. (2004). Let $\tilde{P}_{(1)j} \leq \cdots \leq \tilde{P}_{(q)j}$ be the ordered versions of \tilde{P}_{ij} , $i = 1, \cdots, q$, for a fixed $j = 1, \ldots, m$. Then, in the Bonferroni test, the pooled (or adjusted) p-value is given by $P_j = q\tilde{P}_{(1)j}$; whereas, in the Simes test, it is given by $P_j = \min_{1 \leq i \leq q} \left\{q\tilde{P}_{(i)j}/i\right\}$. While the Bonferroni test does not require any dependence structure in the underlying pvalues, the Simes test requires a certain type of positive dependence condition that is often satisfied in multiple testing applications (Sarkar and Chang, 1997). Upon rejection of H_{0j} using the Bonferroni pooled p-value at a level α , the *i*th component null hypothesis H_{0j}^i can be rejected if $\tilde{P}_{ij} \leq \alpha/q$. For the test based on the Simes pooled p-value, H_{0j}^i corresponding to every $\tilde{P}_{ij} \leq \tilde{P}_{(R_j)j}$ is rejected, where $R_j = \max\left\{i: \tilde{P}_{(i)j} \leq \frac{i}{q}\alpha\right\}$, if the maximum exists; otherwise, $R_j = 0$.

Now, suppose the pooled *p*-value P_j , based on either Bonferroni or Simes test, is available to us for every j = 1, ..., m, to carry out a multiple testing procedure to test H_{0j} against H_{1j} simultaneously for all j = 1, ..., m. We will use the multiple testing method of Benjamini and Hochberg (1995) (the BH method) that is designed to control the false discovery rate (FDR). The FDR, for any given multiple testing procedure, is the expected proportion of false rejections (Type I errors) among all rejections, an overall measure of Type I error rate commonly used in microarray studies. More formally, with V the number of falsely rejected true null hypotheses among H_1, \ldots, H_m and R the total number of rejected hypotheses among H_1, \ldots, H_m , it is defined as

$$FDR = E\left\{\frac{V}{R \vee 1}\right\},\tag{3}$$

where $R \vee 1 = \max(R, 1)$. This method with a control of the FDR at a given level α is a stepup test that, given ordered *p*-values $P_{(1)} \leq \cdots \leq P_{(m)}$ with the corresponding null hypotheses $H_{(1)}, \cdots, H_{(m)}$, finds $k = \max \left\{ 1 \leq j \leq m : P_{(j)} \leq j\alpha/m \right\}$ and rejects those $H_{(j)}$ for which $P_{(j)} \leq P_{(k)}$, provided this maximum exists, otherwise, accepts all the null hypotheses.

When a H_{0j} : $\delta_j = 0$ is rejected using the BH method and further decisions are being made on the signs of the component δ_{ij} 's in the corresponding δ_j , a directional error might occur due to wrong assignments of the signs. For instance, if there is a component δ_{ij} in $\delta_j = (\delta_{1j}, \ldots, \delta_{qj})$ that is truly positive (or negative) but declared to be negative (or positive) while deciding on the signs of the δ_{ij} 's upon rejection of H_{0j} : $\delta_j = 0$, a directional error occurs. So, we need to control such directional errors as well. A convenient and practical way of doing that would be to use an error rate combining both Type I and directional errors in the FDR framework and make sure that it is controlled. One such error rate is the mixed directional FDR (mdFDR), the sum of the FDR and the pure directional FDR (dFDR). The dFDR is defined as

$$dFDR = E\left\{\frac{S}{R \vee 1}\right\},\tag{4}$$

where S denotes the total number of false null hypotheses among H_1, \ldots, H_m that are correctly rejected but at least one directional error has been made while deciding upon the signs of the components. In other words, S is the number of rejected hypotheses H_j 's such that $\delta_j \neq 0$ and for some $i = 1, \ldots, q$, δ_{ij} is declared to be positive when $\delta_{ij} \leq 0$, or δ_{ij} is declared to be negative when $\delta_{ij} \geq 0$. Thus, more formally, the mdFDR is defined as

$$mdFDR = FDR + dFDR = E\left\{\frac{V+S}{R\vee 1}\right\},\tag{5}$$

the expected proportion of Type I and directional errors among all rejections.

It is important to point out that the goal of this paper is to identify expression patterns of m genes over p ordered categories. For each gene it is biologically relevant to consider its expression pattern as a whole across p ordered categories rather than viewing this to be a problem of testing qm separate hypotheses which ignores the intrinsic biological structure present in the problem. Thus, rather than viewing it as a problem of performing qm tests, we treat it as a problem of performing a set of m tests each involving q-dimensional hypothesis. In addition, we want to emphasize that while making directional decisions for the components of δ_j , no directional errors are being made when $\delta_j = 0$. In contrast, when making directional decisions regarding a non-null δ_j , a directional error is made if a component δ_{ij} for which $\delta_{ij} = 0$ is declared to be positive or negative.

In the next section, we will develop methods to control the mdFDR. This will extend the following directional BH procedure of Benjamini and Yekutieli (2005) from dimension one (i.e., q = 1) to a general dimension.

DEFINITION 1 (The level- α directional BH Procedure)

- (1) Apply the BH method at level α to test $H_{0j} : \delta_{1j} = 0$ against $H_{1j} : \delta_{1j} \neq 0$ simultaneously for j = 1, ..., m, based on the two-sided p-values $\tilde{P}_{1j}, j = 1, ..., m$.
- (2) Let R denote the total number of null hypotheses rejected.
- (3) For every $j = 1, \dots, m$, with $\tilde{P}_{1j} \leq \frac{R}{m} \alpha$, declare $\delta_{1j} > or < 0$ according as $T_{1j} > 0$ or < 0.

It controls the mdFDR at α under independence of the underlying test statistics.

3. Multidimensional Directional FDR Controlling Procedures

We introduce in this section our proposed method of controlling the mdFDR while testing $H_{0j}: \delta_j = \mathbf{0}$ against $H_{1j}: \delta_j \neq \mathbf{0}$, simultaneously for all j = 1, ..., m, and making further decisions on the signs of the δ_{ij} 's upon rejection of the corresponding H_{0j} . It is based on the Bonferroni pooled *p*-values.

Procedure 1

- Apply the BH method at level α to test H_{0j} against H_{1j} simultaneously for j = 1,...,m,
 based on the Bonferroni pooled p-values P_j, j = 1,...,m.
- (2) Let R denote the total number of null hypotheses rejected.
- (3) For every $i = 1, \dots, q$ and $j = 1, \dots, m$ with $\tilde{P}_{ij} \leq \frac{R}{qm} \alpha$, if $T_{ij} > 0$, declare $\delta_{ij} > or < 0$ according as $T_{ij} > 0$ or < 0.

THEOREM 1: With independent q-dimensional test statistics $\mathbf{T}_j = (T_{1j}, \dots, T_{qj}), j = 1, \dots, m$, the mdFDR of Procedure 1 is less than or equal to α .

Remark 1. We offer a proof of Theorem 1 in the Appendix. Benjamini and Yekutieli (2005) gave an indirect proof of this theorem in the special case when q = 1 using an approach that relates to the FDR-adjusted confidence intervals for selected parameters they developed in the same paper. However it is not apparent how one could adapt their proof to the present case involving multiple parameters. So, we provide a direct proof in a more general setting.

Remark 2. In Theorem 1, we assume that q-dimensional test statistics \mathbf{T}_j 's are independent. However, within each \mathbf{T}_j , we do not impose any restriction on T_{ij} 's.

It would be tempting to develop an alternative method based on the Simes pooled *p*-values as follows:

Procedure 2

- Apply the BH method at level α to test H_{0j} against H_{1j} simultaneously for j = 1,..., m, based on the Simes pooled p-values P_j, j = 1,..., m.
- (2) Let R denote the total number of null hypotheses rejected.
- (3) For every $j = 1, \dots, m$, let $\tilde{P}_{(1)j} \leq \dots \leq \tilde{P}_{(q)j}$ be the ordered values of $\tilde{P}_{ij}, i = 1, \dots, q$.

Let $R_j = \max\{i : \tilde{P}_{(i)j} \leq \frac{i}{q} \cdot \frac{R}{m}\alpha\}$, if the maximum exists; otherwise $R_j = 0$. For every iand j with $\tilde{P}_{ij} \leq \frac{R_j}{q} \cdot \frac{R}{m}\alpha$, declare $\delta_{ij} > 0$ or < 0 according as $T_{ij} > 0$ or < 0.

Remark 3. As Simes test is known to be more powerful than the Bonferroni test (Simes, 1986), Procedure 2 would be more powerful than Procedure 1. Unfortunately, however, it would not control the mdFDR, as the Associate Editor pointed out. Consider, for instance, m = 1. The augmented test in this procedure in this case reduces to the step-up test with Simes critical values for the q hypotheses. Assume that q = 10 and that for half of the hypotheses $\delta_{i1} = 0$ and for the remaining δ_{i1} is very large. Then the familywise error rate (FWER) of the step-up test with Simes critical values (for the test of the q hypotheses) is not controlled; see also Hommel (1988). However, in this scenario, mdFDR \geq FWER. Therefore, Procedure 2 loses the control of the mdFDR in this situation. So, we do not formally propose it in this article as a multidimensional directional FDR controlling procedure, though we will consider it along with Procedure 1 in our simulation studies in the next section.

4. A Simulation Study

A simulation study was performed to evaluate the performance of our proposed method, Procedure 1. Specifically, we wanted to investigate the following three questions: (i) How does it perform in terms of its control of the FDR, dFDR and mdFDR and also power under independence as well as under types of dependence across the genes? (ii) How dose it perform in terms of the same operating characteristics under the independence across the genes when we benchmark it against Procedure 2 (based on the Simes

pooled *p*-values) and the procedure that makes no adjustment to the gene specific *p*-values, that is, simply uses $\tilde{P}_{(1)j}$ as the pooled *p*-value?

(iii) How does the performance of Procedure 1 under the independence across the genes change as the the dimension q increases?

We generated q + 1 independently distributed *m*-dimensional random normal vectors $\mathbf{Z}_1, \ldots, \mathbf{Z}_{q+1}$, where the components $Z_{ij}, j = 1, \cdots, m$, in each \mathbf{Z}_i are dependent with $Z_{ij} \sim$ $N(\mu_{ij},1)$ and have a common correlation ρ . Let $\delta_{ij} = (\mu_{i+1,j} - \mu_{ij})/\sqrt{2}, i = 1, \ldots, q; j =$ $1, \ldots, m$. Out of the *m* parameter vectors $\boldsymbol{\delta}_j = (\delta_{1j}, \ldots, \delta_{qj}), j = 1, \ldots, m, m_0$ were set to a null vector each, and all the δ_{ij} 's in 50%, 25% and 25% of the remaining $m-m_0$ δ_j 's were selected randomly from the intervals (-0.75, 0.75), (-4.25, -2.75) and (2.75, 4.25) respectively. For each $i = 1, \dots, q$, and $j = 1, \dots, m$, the statistic $T_{ij} = (Z_{i+1,j} - Z_{ij})/\sqrt{2}$ for testing H_{0j}^i : $\delta_{ij} = 0$ vs. H_{1j}^i : $\delta_{ij} \neq 0$ and the corresponding two-sided *p*-value $\tilde{P}_{ij} = 2\{1 - \Phi(|T_{ij}|)\}$ were then computed, where $\Phi(\cdot)$ is the standard normal cdf. The pooled *p*-values were calculated according to the Bonferroni and Simes tests, respectively, and Procedures 1, 2 were applied to their respective lists of pooled p-values for testing the mnull hypotheses described in (1). We also considered the so called no-adjustment procedure, which are the same as Procedures 1 and 2, except for every hypothesis H_j we do not make any adjustment for its corresponding p-value $P_j = \tilde{P}_{(1)j}$. For each of these procedures, the number of true null hypotheses that are rejected (Type I errors), the number of δ_i 's corresponding to the false null hypotheses the signs of whose components do not completely match with those assigned by the procedure (directional errors) and the sum of these two numbers (Type I and directional errors) were noted. Finally, the following three proportions among the total number of rejected null hypotheses were calculated – the proportion of Type I errors (the observed value of $V/R \vee 1$), the proportion of directional errors (the observed value of $S/R \vee 1$) and the proportion of Type I and directional errors (the observed value of $(V + S)/R \vee 1$). These steps were repeated 10,000 times and the simulated values of the FDR, dFDR and mdFDR were obtained by averaging out the 10,000 values of the above three proportions.

[Figure 1 about here.]

[Figure 2 about here.]

Figure 1 presents the simulated FDR, dFDR and mdFDR, Figure 2 presents standard deviation of the simulated mdFDR, and Web Figure 1 presents the simulated average power (the proportion of false null hypotheses that are correctly rejected with correct assigned signs) of Procedure 1 plotted against the number of false null hypotheses for m = 1000, q = 5, $\alpha = 0.05$ and $\rho = 0$ (independence), 0.2, 0.5 and 0.8.

Some interesting observations can be made from Figure 1. With increasing number of truly significant genes, the FDR steadily decreases to zero as long as the dependence across the genes is low or moderately high, while the dFDR slowly increases from zero to a value slightly less than 0.01, no matter what the dependence across the genes is, as long as it is non-negative. Consequently, when the genes are not too highly dependent, with increasing number of truly significant genes although the mdFDR decreases, implying that Procedure 1 as an mdFDR controlling procedure becomes more conservative, it does not however reach zero (see Figure 1(a)-(c)). When the genes are highly dependent, as we see from Figure 1(d), Procedure 1 becomes less conservative as the number of truly significant genes begins to increase from zero, but eventually it becomes more conservative as this number becomes larger.

Also, as seen from Figure 2, the standard deviation of the estimated mdFDR is very small. From Web Figure 1 we see that as the dependence across genes increases, the change in power is small. Overall, the effect of dependence across genes on the performance of the proposed procedure is relatively small. As suggested by the associate editor, under the above simulation settings, we also evaluated the performance of Procedure 1 for the $\delta_j = (100, 0, \dots, 0)$, in which one component is extremely large and the rest are zero. For such non-null δ_j , there is a high chance that it is detected to be non-null and its one or more zero components are declared to be positive or negative. That is, in such scenario, it is more possible to make directional errors. Web Figure 2 presents the simulated FDR, dFDR and mdFDR and Web Figure 3 presents the standard deviation of the simulated mdFDR. As expected, in the case of a non-null pattern, the dFDR is increasing as the number of false nulls increases, and is much larger than that for the previous scenario. However, the mdFDR is still controlled under a pre-specified level (see Web Figure 2). As seen from Web Figure 3, the standard deviation of the estimated mdFDR is still very small for such scenarios.

Web Figure 4 presents an answer to question (ii). As we see from this figure, Procedures 1 and 2 behave quite similarly, at least when the dependence across the genes is not of concern, in terms of controlling the FDR, dFDR and mdFDR and the power, though Procedure 2 is slightly more liberal as expected. Also as expected, if no adjustment is made to gene-specific p-values, we lose the control of the FDR and mdFDR, with the maximum reaching 0.2. It seems surprising that, even without any adjustment to gene specific p-values, the dFDR always remains low, though it becomes larger compared to that for Procedures 1 and 2 as the number of false nulls increases.

Web Figures 5-6 provide an answer to question (iii). It is interesting to note that the performance of Procedure 1 in terms of controlling the FDR, dFDR and mdFDR is unaffected by the dimension q when the dependence across the genes is not present. The power, of course, increases with increasing dimension.

5. An Application to Time-Course Gene Expression Data

Lobenhofer et al. (2002) investigated the effect of estrogen on the expression of cell-cycle genes as MCF-7 breast cancer cells go through the cell division cycle. A normal cell division cycle consists of four major phases, namely, the G1 (or Gap 1), S (Synthesis), G2 (or Gap 2) and M (Mitosis) phase. Genes involved in the cell cycle (known as *cell-cycle genes*) are expected to attain peak gene expression during the phase in which they have a specific biological function in the cell cycle.

According to Lobenhofer et al. (2002), most estradiol treated MCF-7 cells are expected

to go through S, G2 and M phases in 12 to 36 hours after treatment and complete the cycle in 48 hours. Genes involved in cell growth and related activities are expected to have maximum expression (or minimum expression if they are anti-growth) during 1 or 4 hours and then monotonically decrease (or increase) in expression as cells go through the remaining phases. On the other hand, genes involved in DNA synthesis, repair and mitosis would have maximum (or minimum) expression during 12 to 36 hours. Thus, such genes may have an *Umbrella* (or *Inverted umbrella*) shaped pattern with a peak or trough during 12 to 36 hours time period. However, according to Lobenhofer et al. (2002), the cells may be asynchronous as they complete the cell division cycle at 48 hours after exposure. For this reason, the expression of some of the cell-cycle genes may not return to their baseline values at 48 hours but may attain a plateau.

Before exposing the MCF-7 breast cancer cells to estrogen, Lobenhofer et al. (2002) first synchronized all the cells to G1 phase by depriving the cells of serum for 24 hours. Synchronization of cells to the same phase at the beginning of the experiment is important for obtaining reliable gene expression data. They then harvested estradiol treated cells after 1, 4, 12, 24, 36 or 48 hours of treatment. Gene expressions using cDNA microarray chips were obtained at each time point. Each cDNA microarray chip consisted of 1900 gene probes. With 8 replicates at each time point, there were a total of 48 microarray chips across the 6 time points.

Motivated by the above observations, in this section we apply the proposed methodology to identify some cell-cycle genes by considering four ordered categories of time points, namely, 1 hour (T1), 4 hours (T2), mid group (i.e., the union of 12, 24, 36 hours) (T3) and 48 hours (T4) after treatment. Thus the sample sizes in the four groups are 8, 8, 24 and 8 respectively. Since the major cell division related activity takes place during the 12 to 36 hours time interval, we combined those time periods together to contrast that period from initial cell growth period (1, 4 hours) and the end of mitosis (48 hours).

Suppose $\theta_{T1,j}$, $\theta_{T2,j}$, $\theta_{T3,j}$ and $\theta_{T4,j}$ denote the mean gene expression of gene j, j = 1, 2, ..., 1900, during time periods T1, T2, T3 and T4, respectively. Using notations from the previous section, we let $\delta_{1j} = \theta_{T1,j} - \theta_{T2,j}$, $\delta_{2j} = \theta_{T2,j} - \theta_{T3,j}$ and $\delta_{3j} = \theta_{T3,j} - \theta_{T4,j}$.

Let T_{ij} denote the test statistic associated with the parameter δ_{ij} , i = 1, 2, 3 and j = 1, 2, ..., 1900. In this application T_{ij} is the usual two-sample *t*-test statistic and since the underlying data are not necessarily normally distributed, nonparametric bootstrap methodology based on 10,000 bootstrap samples is used for computing the *p*-values \tilde{P}_{ij} associated with hypotheses on δ_{ij} . We then calculate the Bonferroni pooled *p*-value P_j for each gene *j* after computing \tilde{P}_{ij} , i = 1, 2, 3.

By applying our proposed method, Procedure 1 to the list of the pooled *p*-values P_j 's, we identified 86 differentially expressed genes at level $\alpha = 0.05$ of which 19 had an umbrella shape response, 3 inverted umbrella, 32 increased in expression from T1 to T3 and then plateaued (i.e. for some gene j, $\theta_{T1,j} \leq \theta_{T2,j} \leq \theta_{T3,j} = \theta_{T4,j}$, with at least one strict inequality). An opposite response was seen with 22 genes which had decreased expression from T1 to T3 and then plateaued (i.e. for some gene j, $\theta_{T1,j} \geq \theta_{T2,j} \geq \theta_{T3,j} = \theta_{T4,j}$, with at least one strict inequality). We also discovered 10 genes that had a flat expression until T3 and then a decrease in response from T3 to T4.

Comparing our results with those of Lobenhofer et al. (2002) and Peddada et al. (2003), we found that of the 86 genes we identified, 39 were also identified in at least one of the two previous papers. This included 8 of 13 DNA replication/repair genes identified by Lobenhofer et al. (2002). Among the 5 that were not identified by our procedure, we note that except for MCM7, which may be significant at $\alpha = 0.10$, all others had large *p*-values that were not significant even at $\alpha = 0.20$. Interestingly, in addition to MCM3 that was identified by both Lobenhofer et al. (2002) and Peddada et al. (2003), we identified a well known cell-cycle gene MCM4 (http://www.cyclebase.org).

An important step in DNA synthesis during the S phase is the binding of complex proteins to DNA for recruiting other proteins necessary for DNA synthesis. One such complex protein is the replication factor C. Lobenhofer et al. (2002) identified one subunit of this protein, known as replication factor C3. Later the order-restricted inference based methodology of Peddada et al. (2003) identified two additional subunits of this protein, namely, replication factors C4 and C5. Interestingly, the newly introduced methodology identified subunits C2, C3 and C5 as significant genes, thus reinforcing the earlier findings and adding one more subunit to the previous list of replication factor C. Furthermore, based on the proposed methodology it is possible to conclude that the subunits C2, C3 and C5 have peak expression during the 12, 24 or 36 hours time period where the DNA synthesis and replication takes place.

Furthermore, similar to the order-restricted inference procedure of Peddada et al. (2003), the proposed methodology identified the cyclin-dependent kinase inhibitor 1 A (p21 and Cip 1) as repressed during 12 to 36 hours. This gene was not identified by Lobenhofer et al. (2002).

A complete list of all genes identified by this procedure is provided in the Web Supplementary Materials accompanying this article.

6. Concluding Remarks

In microarray gene expression studies, researchers are often not only interested in identifying differentially expressed genes under different biological conditions, but are also interested in detecting trends in mean response over ordered categories. For instance, in the simple case of two categories (normal versus tumor tissue), researchers are not only interested in identifying significant genes across these two categories, but they are also interested in further identifying

the down and up-regulated genes. As the number of ordered categories increases, the trends or directional patterns become complex and the number of directional patterns increases. Except for the usual Type I errors, this also potentially results in a relatively high frequency of directional errors. Hence, it is important to develop statistical methods of identifying trends in mean response over ordered categories while maintaining a control over both the Type I and directional errors.

The approach proposed in this article provides such a methodology. Differently from existing statistical methods (Peddada et al., 2003; Lin et al., 2007), we have formulated the problem of identifying trends in mean response over ordered categories as a multiple testing problem involving successive comparisons and further directional decisions on the multidimensional parameter of each gene. To deal with this problem, we have first suggested a general multidimensional BH-type directional procedure using the Bonferroni test for controlling the mixed directional FDR (mdFDR), an overall measure of both Type I and directional errors within the framework of the FDR, and theoretically proved that the proposed procedure controls the mdFDR at a pre-specified level when the underlying test statistics are independent across the genes. We evaluated the performance of the introduced procedure in the case of dependence through a simulation study. Finally, the whole proposed methodology has been applied to analyze a time-course microarray data and some interesting results have been obtained.

Although our focus was on identifying individual gene expression profile or trend over the ordered categories in some common microarray experiments such as time-course or dose-response experiments, the proposed methodology can be also applied in National Toxicology Program (NTP) studies, where researchers are interested in determining whether for a given tumor type, there is a significant dose effect and then identifying its dose-response profile.

The methodology proposed in this article provides an interesting starting point towards ad-

dressing the complex yet important problem of controlling both Type I and directional errors in multiple testing involving multidimensional parameters. The mdFDR controlling property of the proposed directional BH procedure has been established under the assumption that the underlying test statistics are independent across the genes. When gene expressions are obtained by drawing samples from same subjects over time, such an assumption need not be valid. In such cases, not only do we have dependence among gene expressions at a given time point but there may be temporal dependence among gene expressions at different time points. It will be interesting to theoretically investigate the performance of the proposed directional BH procedure under such complex dependence structures. In addition, it will also be interesting to develop more powerful adaptive BH directional FDR procedure by exploiting knowledge of the proportion of true null hypotheses.

7. Supplementary Materials

Web Appendices and Figures referenced in Sections 3 and 4 are available under the Paper Information link at the *Biometrics* website http://www.biometrics.tib.org. A complete list of all genes identified in Section 5 are also available at the Biometrics website.

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LIST OF FIGURES

1 Performance of Procedure 1 under dependence across genes in terms of its control of the FDR, dFDR and mdFDR for $m = 1000, q = 5, \alpha = 0.05$, and $\rho = 0, 0.2, 05$ and 0.8.

2 Standard deviation of the mdFDR of Procedure 1 under dependence across genes for $m = 1000, q = 5, \alpha = 0.05$, and $\rho = 0, 0.2, 05$ and 0.8.



Figure 1. Performance of Procedure 1 under dependence across genes in terms of its control of the FDR, dFDR and mdFDR for $m = 1000, q = 5, \alpha = 0.05$, and $\rho = 0, 0.2, 05$ and 0.8.



Figure 2. Standard deviation of the mdFDR of Procedure 1 under dependence across genes for $m = 1000, q = 5, \alpha = 0.05$, and $\rho = 0, 0.2, 05$ and 0.8.