FORMALIZED DATA SNOOPING BASED ON GENERALIZED ERROR RATES

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Abstract

It is common in econometric applications that several hypothesis tests are carried out at the same time. The problem then becomes how to decide which hypotheses to reject, accounting for the multitude of tests. The classical approach is to control the familywise error rate (FWE), that is, the probability of one or more false rejections. But when the number of hypotheses under consideration is large, control of the FWE can become too demanding. As a result, the number of false hypotheses rejected may be small or even zero. This suggests replacing control of the FWE by a more liberal measure. To this end, we review a number of proposals from the statistical literature. We briefly discuss how these procedures apply to the general problem of model selection. A simulation study and two empirical applications illustrate the methods.

KEY WORDS: Data snooping, false discovery proportion, false discovery rate, generalized familywise error rate, model selection, multiple testing, stepwise methods.

JEL CLASSIFICATION NOS: C12, C14, C52.
1 Introduction

Much empirical research in economics and finance inevitably involves data snooping. The problem arises when several hypothesis tests are carried out at once and one has to decide which hypotheses to reject. One common scenario is the comparison of many strategies (such as investment strategies) to a common benchmark (such as a market index); here, a particular hypothesis test specifies whether a particular strategy outperforms the benchmark or not. Another common scenario is multiple regression models; here, a particular hypothesis test specifies whether a particular regression coefficient is equal to a prespecified value or not.

Economists have long been aware of the dangers resulting from data snooping; see, for example, White (2000), Hansen (2005), Romano and Wolf (2005c), and the references therein. The standard approach to account for data snooping is to control (asymptotically) the familywise error rate (FWE), which is the probability of making one or more false rejections; e.g., see Westfall and Young (1993). However, this criterion can be too strict when the number of hypotheses under consideration is very large. As a result, it can become very difficult (or impossible) to make true rejections. In other words, controlling the FWE can be 'playing it too safe'.

When the number of hypotheses is very large and the ability to make true rejections is a main concern, it has been suggested to relax control of the FWE. In this paper, we discuss and review three proposals to this end. The first proposal is to control the probability of making $k$ or more false rejections, for some integer $k$ greater than or equal to one, which is called the $k$-FWE. The remaining proposals are based on the false discovery proportion (FDP), defined as the number of false rejections divided by the total number of rejections; and defined to be 0 if there are no rejections at all. The second proposal is to control $E(\text{FDP})$, the expected value of the FDP, which is called the false discovery rate (FDR). The third proposal is to control $P\{\text{FDP} > \gamma\}$, where $\gamma$ is a small, user-defined number; the special case $\alpha = 0.5$ yields control of the median FDP. While the three proposals are different, they share a common philosophy: by allowing a small number or a small (expected) proportion of false rejections, one can improve one's chances to make true rejections, and perhaps greatly so. In other words, the price to pay can be small compared to the benefits to reap.

This paper reviews various methods that have been suggested for control of the three criteria previously mentioned, including some very recent multiple testing procedures that account for the dependence structure of the individual test statistics. Part of our contribution is to present the

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1 We use the compact terminology of false rejection to denote the rejection of a true null hypothesis. Similarly, the terminology true rejection denotes the rejection of a false null hypothesis. A false rejection is sometimes coined a false discovery.

2 The 1-FWE is simply the usual FWE.
methods in a unified context, allowing an applied researcher to grasp the concepts quickly, rather than having to read and digest the numerous underlying original papers. We also demonstrate, by means of some simulations and two empirical applications, how competing multiple testing procedures compare when applied to data.

The remainder of the paper is organized as follows. Section 2 describes the model and the formal inference problem. Section 3 reviews various methods to control the FWE. Section 4 presents various methods to control the k-FWE. Section 5 reviews the method of Benjamini and Hochberg (1995) for control of the FDR. Section 6 presents various methods to control \( P\{\text{FDP} > \gamma \} \). Section 7 discusses applications of generalized error rates to the problem of model selection. Section 8 sheds some light on finite-sample performance of the discussed methods via a simulation study. Section 9 provides two empirical applications. Finally, Section 10 concludes. An appendix contains some details concerning bootstrap implementation.

2 Notation and Problem Formulation

One observes a data matrix \( x_{t,l} \) with \( 1 \leq t \leq T \) and \( 1 \leq l \leq L \). The data is generated from some underlying probability mechanism \( P \) which is unknown. The row index \( t \) corresponds to distinct observations, and there are \( T \) of them. In our asymptotic framework, \( T \) will tend to infinity. The column index \( L \) is fixed. For compactness, we introduce the following notation: \( X_T \) denotes the complete \( T \times L \) data matrix; \( X_{t,.}^{(T)} \) is the \( L \times 1 \) vector that corresponds to the \( t \)th row of \( X_T \); and \( X_{.,l}^{(T)} \) is the \( T \times 1 \) vector that corresponds to the \( l \)th column of \( X_T \).

Interest focuses on a parameter vector \( \theta = \theta(P) \) of dimension \( S \), that is, \( \theta = (\theta_1, \ldots, \theta_S)' \). The individual hypotheses are about the elements of \( \theta \) and can be all one-sided and of the form

\[
H_s: \theta_s \leq \theta_{0,s} \quad \text{vs.} \quad H'_s: \theta_s > \theta_{0,s}
\]

or they can be all two-sided and of the form

\[
H_s: \theta_s = \theta_{0,s} \quad \text{vs.} \quad H'_s: \theta_s \neq \theta_{0,s}.
\]

For each hypothesis \( H_s, 1 \leq s \leq S \), one computes a test statistic \( w_{T,s} \) from the data matrix \( X_T \). In some instances, we will also consider studentized test statistics \( z_{T,s} = w_{T,s} / \hat{\sigma}_{T,s} \), where the standard error \( \hat{\sigma}_{T,s} \) estimates the standard deviation of \( w_{T,s} \) and is also computed from \( X_T \). In the sequel, we often call \( w_{T,s} \) a ‘basic’ test statistic to distinguish it from the studentized statistic \( z_{T,s} \).

We now introduce some compact notation: the \( S \times 1 \) vector \( W_T \) collects the individual basic test
statistics $w_{T,s}$; the $S \times 1$ vector $Z_T$ collects the individual studentized test statistics $z_{T,s}$.

A multiple testing method yields a decision concerning each individual testing problem by either rejecting $H_s$ or not. In an ideal world, one would like to reject all those hypotheses that are false. In a realistic world, and given a finite amount of data, this cannot be achieved with certainty. At this point, we vaguely define our goal as making as many true rejections as possible while not making 'too many' false rejections. Different notions of accounting for data snooping entertain different views of what constitutes 'too many' false rejections.

We next describe two broad examples where data snooping arises naturally and by putting them in the above framework.

**Example 2.1 (Comparing Several Strategies to a Common Benchmark)** One considers $S$ strategies (such as investment strategies) and compares them to a common benchmark (such as a market index). The data matrix $X_T$ has $L = S + 1$ columns then: the first $S$ columns record the individual strategies while the last column records the benchmark. The goal is to decide which strategies outperform the benchmark. Here the individual parameters are defined in such a way that $\theta_s \leq 0$ if and only if the $s$th strategy does not outperform the benchmark. One then is in the one-sided setup (1) with $\theta_{0,s} = 0$ for $s = 1, \ldots, S$. ■

To make this paper self-contained, we now repeat some specific scenarios where several strategies are compared to a benchmark, together with the corresponding test statistics, from Romano and Wolf (2005c).

**Example 2.1.(a) (Absolute Performance of Investment Strategies)** Historical returns of investment strategy $s$, say a particular mutual fund or a particular trading strategy, are recorded in $X_{-s}^{(T)}$. Historical returns of a benchmark, say a stock index or a buy-and-hold strategy, are recorded in $X_{S+1}^{(T)}$. Depending on preference, these can be 'real' returns or log returns; also, returns may be recorded in excess of the risk free rate if desired. Let $\mu_s$ denote the population mean of the return for strategy $s$. Based on an absolute criterion, strategy $s$ beats the benchmark if $\mu_s > \mu_{S+1}$. Therefore, we define $\bar{\theta}_s = \mu_s - \mu_{S+1}$. Using the notation

$$\bar{x}_{T,s} = \frac{1}{T} \sum_{t=1}^{T} x_{t,s}$$

a natural basic test statistic is

$$w_{T,s} = \bar{x}_{T,s} - \bar{x}_{T,S+1} . \tag{3}$$
Often, a studentized statistic is preferable and is given by

\[ z_{T,s} = \frac{\bar{x}_{T,s} - \bar{x}_{T,S+1}}{\hat{\sigma}_{T,s}}, \]  

(4)

where \( \hat{\sigma}_{T,s} \) is an estimator of the standard deviation of \( \bar{x}_{T,s} - \bar{x}_{T,S+1} \).

**Example 2.1.(b) (Relative Performance of Investment Strategies)** The basic setup is as in the previous example, but now consider a risk-adjusted comparison of the investment strategies, based on the respective Sharpe ratios. With \( \mu_s \) again denoting the mean of the return of strategy \( s \) and with \( \sigma_s \) denoting its standard deviation, the corresponding Sharpe ratio is defined as \( SR_s = \frac{\mu_s}{\sigma_s}. \) An investment strategy is now said to outperform the benchmark if its Sharpe ratio is higher than the one of the benchmark. Therefore, we define \( \theta_s = SR_s - SR_{S+1} \). Let

\[ s_{T,s} = \sqrt{\frac{1}{T-1} \sum_{t=1}^{T} (x_{t,s} - \bar{x}_{T,s})^2}. \]

Then a natural basic test statistic is

\[ w_{T,s} = \frac{\bar{x}_{T,s} - \bar{x}_{T,S+1}}{s_{T,s}}. \]  

(5)

Again, a preferred statistic might be obtained by dividing by an estimate of the standard deviation of this difference.

**Example 2.1.(c) (CAPM alpha)** Historical returns of investment strategy \( s \), in excess of the risk-free rate, are recorded in \( X_{t,s}^{(T)} \). Historical returns of a market proxy, in excess of the risk-free rate, are recorded in \( X_{s,S+1}^{(T)} \). For each strategy \( s \), a simple time series regression

\[ x_{t,s} = \alpha_s + \beta_s x_{t,S+1} + \epsilon_{t,s}, \]  

(6)

is estimated by ordinary least squares (OLS). If the Capital Asset Pricing Model (CAPM) holds, all intercepts \( \alpha_s \) are equal to zero. So the parameter of interest here is \( \theta_s = \alpha_s \). Since the CAPM may be violated in practice, a financial advisor might want to identify investment strategies which have a positive \( \alpha_s \). Hence, an obvious basic test statistic would be

\[ w_{T,s} = \hat{\alpha}_{T,s}. \]  

(7)

---

\(^3\)The definition of a Sharpe ratio is often based on returns in excess of the risk-free rate. But for certain applications, such as long-short investment strategies, it can be more suitable to base it on the nominal returns.

\(^4\)We trust there is no possible confusion between a CAPM alpha \( \alpha_s \) and the level \( \alpha \) of multiple testing methods discussed later on.
Again, it can be advantageous to studentize by dividing by \( \hat{\sigma}_{T,s} \), an estimated standard deviation of \( \hat{\sigma}_{T,s} \).

**Example 2.2 (Multiple Regression)** Consider the multiple regression model

\[
y_t = \theta_1 x_{1,t} + \ldots + \theta_H x_{H,t} + \epsilon_t \quad t = 1, \ldots, T.
\]

The data matrix \( X_T \) has \( L = H + 1 \) columns: the first \( H \) columns record the explanatory variables while the last column records the response variable, letting \( x_{H+1,t} = y_t \). Of interest are \( S \leq H \) of the regression coefficients. Without loss of generality, assume that the explanatory variables are ordered in such a way that the coefficients of interest correspond to the first \( S \) coefficients, so \( \theta = (\theta_1, \ldots, \theta_S)' \). One typically is in the two-sided setup (2) where the prespecified values \( \theta_{0,s} \) depend on the context, but at times the one-sided setup (1) can be more appropriate.

In much applied research, all the regression coefficients are of interest—except possibly an intercept if it is included in the regression—and one would like to decide which of them are different from zero. This corresponds to the two-sided setup (2) where \( S = H \)—or \( S = H - 1 \) in case of an included intercept whose coefficient is not of interest—and \( \theta_{0,s} = 0 \) for \( s = 1, \ldots, S \).

Let \( \hat{\theta}_T \) denote an estimator of \( \theta \) computed from the data matrix \( X_T \), using ordinary least squares (OLS) or feasible generalized least squares (FGLS), say. Then the ‘basic’ test statistic for \( H_S \) is simply \( w_{T,s} = \hat{\theta}_{T,s} \). The proper choice of the standard error \( \hat{\sigma}_{T,s} \) for studentization depends on the context. In the simplest case, it can be the usual OLS standard error. More generally, a standard error which is robust against heteroskedasticity and/or autocorrelation might be required; for example, see White (2001).

For testing an individual hypothesis \( H_s \) based on a studentized test statistic \( z_{T,s} \), one can typically compute an approximate \( p \)-value by invoking asymptotic standard normality. For example, for testing a one-sided hypothesis \( H_s : \theta_s \leq \theta_{0,s} \), one might compute \( \hat{p}_{T,s} = 1 - \Phi(z_{T,s}) \), where \( \Phi(\cdot) \) is the standard normal c.d.f.. Of course, one may also appeal to other techniques which rely on approximating the null distribution of \( z_{T,s} \), such as bootstrapping, subsampling, permutation tests, empirical likelihood, or Edgeworth approximations. In any case, \( \hat{p}_{T,s} \) is a marginal \( p \)-value in the sense that the test which rejects \( H_s \) if \( \hat{p}_{T,s} \leq \alpha \) has asymptotic rejection probability \( \alpha \) if \( H_s \) is true. It follows that if all \( S \) null hypotheses are true, and we reject \( H_s \) whenever \( \hat{p}_{T,s} \leq \alpha \), the expected number of false rejections is \( S \times \alpha \) (asymptotically). For example, if \( S = 1,000 \) and \( \alpha = 0.05 \), the expected number of false rejections is 50 (asymptotically) when all null hypotheses are true. Such an approach is too liberal and does not account for the multitude of tests under study. In
the remainder of the paper, we consider various measures of error control which attempt to control false rejections by accounting for the fact that $S$ tests are being carried out simultaneously.

3 Methods Controlling the FWE

The usual approach to deal with data snooping is trying to avoid any false rejections. That is, one seeks to control the familywise error rate (FWE). The FWE is defined as the probability of rejecting at least one of the true null hypotheses. More specifically, if $P$ is the true probability mechanism, let $I_0 = I_0(P) \subset \{1, \ldots, S\}$ denote the indices of the set of true hypotheses, that is,

$$s \in I_0 \text{ if and only if } \begin{cases} \theta_s \leq \theta_{0,s} \text{ in setup (1)} \\ \theta_s = \theta_{0,s} \text{ in setup (2)} \end{cases}.$$ 

The FWE is the probability under $P$ that any $H_s$ with $s \in I_0$ is rejected.\footnote{To show its dependence on $P$, we may write $\text{FWE} = \text{FWE}_P$.}

$$\text{FWE}_P = P\{\text{Reject at least one } H_s : s \in I_0(P)\}.$$ 

In case all the individual null hypotheses are false, the FWE is equal to zero by definition.

Control of the FWE requires that, for any $P$, the FWE be no bigger than $\alpha$, at least asymptotically. Since this must hold for any $P$, it must hold not just when all null hypotheses are true (which is called weak control), but also when some are true and some are false (which is called strong control). As remarked by Dudoit et al. (2003), this distinction is often ignored. The remainder of the paper equates control of the FWE with strong control, and similarly for the control of the $k$-FWE and FDP discussed in later sections. A multiple testing method is said to control the FWE at level $\alpha$ if $\text{FWE}_P \leq \alpha$ for any sample size $T$ and any $P$. A multiple testing method is said to control the FWE asymptotically at level $\alpha$ if $\limsup_{T \to \infty} \text{FWE}_P \leq \alpha$ for any $P$. Methods that control the FWE in finite samples can typically only be derived in special circumstances; see Romano and Wolf (2005b).

3.1 The Bonferroni Method

The most familiar multiple testing method for controlling the FWE is the Bonferroni method. For each null hypothesis $H_s$, one computes an individual $p$-value $\hat{p}_{T,s}$. The Bonferroni method at level $\alpha$ rejects $H_s$ if $\hat{p}_{T,s} \leq \alpha/S$ and therefore is very simple to apply. Because all $p$-values are compared to a single critical value, the Bonferroni method is an example of a single-step procedure. The
disadvantage of the Bonferroni method is that it is in general conservative, resulting in a loss of power.

The Bonferroni method controls the FWE if the distribution of each p-value corresponding to a true null hypothesis is stochastically dominated by the Uniform (0, 1) distribution, that is,

\[ H_s \text{ true } \implies P\{\hat{p}_{T,i} \leq u\} \leq u \quad \text{for any } u \in (0,1) . \]  

(8)

The Bonferroni method asymptotically controls the FWE if the distribution of each p-value corresponding to a true null hypothesis is stochastically dominated by the Uniform (0, 1) distribution asymptotically, that is,

\[ H_s \text{ true } \implies \limsup_{T \to \infty} P\{\hat{p}_{T,i} \leq u\} \leq u \quad \text{for any } u \in (0,1) . \]  

(9)

3.2 The Holm Method

An improvement over Bonferroni is due to Holm (1979) and it works in a stepwise fashion as follows. The individual p-values are ordered from smallest to largest: \( \hat{p}_{T,(1)} \leq \hat{p}_{T,(2)} \leq \ldots \leq \hat{p}_{T,(S)} \) with their corresponding null hypotheses labeled accordingly: \( H_{(1)}, H_{(2)}, \ldots, H_{(S)} \). Then, \( H_{(j)} \) is rejected at level \( \alpha \) if \( \hat{p}_{T,(j)} \leq \alpha/(S - j + 1) \) for \( j = 1, \ldots, S \). In comparison with the Bonferroni method, the criterion for the smallest p-value is equally strict, \( \alpha/S \), but it becomes less and less strict for the larger p-values. Hence, the Holm method will typically reject more hypotheses is more powerful than the Bonferroni method. On the other hand, the Holm method (asymptotically) controls the FWE under exactly the same condition as the Bonferroni method.

The Holm method starts with examining the most significant hypothesis, corresponding to the smallest p-value, and then moves 'down' to the less significant hypothesis. Such stepwise methods are called stepdown methods. Different in nature are stepup methods, which start by examining the least significant hypothesis, corresponding to the largest p-value, and then move 'up' to the more significant hypotheses. An example is the stepwise method of Benjamini and Hochberg (1995); see Section 5.

While its improvement over Bonferroni can be substantial, the Holm method can also be very conservative. The reason for the conservativeness of the Bonferroni and the Holm methods is that they do not take into account the dependence structure of the individual p-values. Loosely speaking, they achieve control of the FWE by assuming a worst-case dependence structure. If the true dependence structure could (asymptotically) be accounted for, one should be able to (asymptotically) control the FWE but at the same time increase power. In many economic or
financial applications, the individual test statistics are jointly dependent. It is therefore important to account for the underlying dependence structure in order to avoid being overly conservative.

3.3 The Bootstrap Reality Check and the StepM Method

White (2000), in the context of Example 2.1, proposes the bootstrap reality check (BRC). The BRC estimates the sampling distribution of $\max_{1 \leq s \leq S}(w_{T,s} - \theta_s)$, implicitly taking into account the dependence structure of the individual test statistics. Let $s_{\text{max}}$ denote the index of the strategy with the largest statistic $w_{T,s}$. The BRC decides whether or not to reject $H_{s_{\text{max}}}$ at level $\alpha$, asymptotically controlling the FWE. It therefore addresses the question whether the strategy that appears 'best' in the observed data really beats the benchmark.\(^6\) On the other hand, it does not attempt to identify as many outperforming strategies as possible.

Hansen (2005) offers some improvements over the BRC. First, his method reduces the influence of ‘irrelevant’ strategies, meaning strategies that ‘significantly’ underperform the benchmark. Second, he proposes the use of studentized test statistics $z_{T,s}$ instead of basic test statistics $w_{T,s}$. However, the method of Hansen (2005) also only addresses the question whether the strategy that appears 'best' in the observed data really beats the benchmark.

Romano and Wolf (2005c), also in the context of Example 2.1, address the problem of detecting as many outperforming strategies as possible. Often, this will be the relevant problem. For example, if a bank wants to invest money in trading strategies that outperform a benchmark, it is preferable to build a portfolio of several strategies rather than fully invest in the ‘best’ strategy only. Hence, the goal is to identify the universe of all outperforming strategies for maximum diversification. The stepwise multiple testing (StepM) method of Romano and Wolf (2005c) improves upon the single-step BRC of White (2000) very much in the way that the stepwise Holm method improves upon the single-step Bonferroni method: in terms of being able to detect more outperforming strategies, one is afforded a free lunch. Like the Holm method, the StepM method is of the stepdown nature; that is, it starts by examining the most significant hypothesis.

While Romano and Wolf (2005c) develop their StepM method in the context of Example 2.1, it is straightforward to adapt it to the generic multiple testing problems (1) and (2). For details, see Subsubsection 4.3.3.

\(^6\)Equivalently, it addresses the question whether there are any strategies at all that beat the benchmark.
4 Methods Controlling the $k$-FWE

By relaxing the strict FWE criterion one will be able to reject more false hypotheses. This section presents the alternative criterion of controlling the $k$-FWE. The $k$-FWE is defined as the probability of rejecting at least $k$ of the true null hypotheses. As before, if $P$ is the true probability mechanism, let $I_0 = I_0(P) \subset \{1, \ldots, S\}$ denote the indices of the set of true hypotheses. The $k$-FWE is the probability under $P$ that any $k$ or more of the $H_s$ with $s \in I_0$ are rejected:

$$k\text{-FWE}_P = P\{\text{Reject at least } k \text{ of the } H_s : s \in I_0\}.$$  

In case at least $S - k + 1$ of the individual null hypotheses are false, the $k$-FWE is equal to zero by definition.

A multiple testing method is said to control the $k$-FWE at level $\alpha$ if $k\text{-FWE}_P \leq \alpha$ for any sample size $T$ and for any $P$. A multiple testing method is said to control the FWE asymptotically at level $\alpha$, if $\limsup_{T \to \infty} k\text{-FWE}_P \leq \alpha$ for any $P$. Methods that control the $k$-FWE in finite samples can typically only be derived in special circumstances; see Romano and Wolf (2005a).

We now describe how the various methods of Section 3 can be generalized to achieve (asymptotic) control of the $k$-FWE. Of course, since our goal is to reject as many false hypotheses as possible, attention will focus on the generalization of the StepM method.

4.1 Generalization of the Bonferroni Method

The generalized Bonferroni method is due to Hommel and Hoffman (1988) and Lehmann and Romano (2005) and is based on the individual $p$-values. The method rejects $H_s$ if $\hat{p}_{T,s} \leq k\alpha/S$. It is easy to see that potentially many more hypotheses will be rejected compared to the original Bonferroni method. Indeed, the cut-off value for the individual $p$-values is $k$ times as large.

If condition (8) holds, then this method controls the $k$-FWE. If condition (9) holds, then this method asymptotically controls the $k$-FWE.

4.2 Generalization of the Holm Method

The individual $p$-values are ordered from smallest to largest, $\hat{p}_{T,(1)} \leq \hat{p}_{T,(2)} \leq \ldots \leq \hat{p}_{T,(S)}$, with their corresponding null hypotheses labeled accordingly, $H_{(1)}$, $H_{(2)}$, $\ldots$, $H_{(S)}$. Then $H_{(s)}$ is rejected
at level $\alpha$ if $p_{T,(j)} \leq \alpha_j$ for $j = 1, \ldots, s$, where\footnote{The $\alpha_j$ depend also on $S$ and $k$, but this dependence is suppressed in the notation.}

$$\alpha_j = \begin{cases} \frac{ka}{S} & \text{for } j \leq k \\ \frac{ka}{S+k-j} & \text{for } j > k \end{cases}$$

This modification is also due to Hommel and Hoffman (1988) and Lehmann and Romano (2005). It is easy to see that this stepwise method is more powerful than the single-step generalized Bonferroni method. On the other hand, the sufficient conditions for control and asymptotic control, respectively, of the $k$-FWE are identical.

## 4.3 Generalization of the StepM Method

We now describe how to modify the StepM method of Romano and Wolf (2005c) in order to achieve asymptotic control of the $k$-FWE. We begin by discussing the one-sided setup (1) and then describe the necessary modifications for the two-sided setup (2).

### 4.3.1 Basic Method

We detail our method in the context of using basic test statistics $w_{T,s}$, and discuss the extension to the studentized case later on. The method begins by relabeling the strategies according to the size of the individual test statistics, from largest to smallest. Label $r_1$ corresponds to the largest test statistic and label $r_s$ to the smallest one, so that $w_{T,r_1} \geq w_{T,r_2} \geq \cdots \geq w_{T,r_s}$.

Some further notation is required. Suppose $\{y_s : s \in K\}$ is a collection of real numbers indexed by a finite set $K$ having $|K|$ elements. Then, for $k \leq |K|$, the $k$-$\max_{s \in K}(y_s)$ is used to denote the $k$th largest value of the $y_s$ with $s \in K$. So, if the elements $y_s$, $s \in K$, are ordered as

$$y(1) \leq \cdots \leq y(|K|) \;,$$

then

$$k$-$\max_{s \in K}(y_s) = y(|K|-k+1) \;.$$ 

Further, for any $K \subset \{1, \ldots, S\}$, define

$$c_K(1 - \alpha, k, P) = \inf \{x : P\{k$-$\max_{s \in K}(w_{T,r_s} - \theta_{r_s}) \leq x\} \geq 1 - \alpha\} \;.$$  \hfill (10)
that is, \( c_K(1 - \alpha, k, P) \) is the smallest \( 1 - \alpha \) quantile of the sampling distribution under \( P \) of 

\[
\text{k-max}_{s \in K}(w_{T,r_s} - \theta_{r_s}).
\]

In the first step of the procedure, we construct a rectangular joint region\(^8\) for the vector 

\[(\theta_{r_1}, \ldots, \theta_{r_S})' \]

of the form 

\[
[w_{T,r_1} - c_1, \infty) \times \ldots \times [w_{T,r_S} - c_1, \infty).
\] (11)

Individual decision are then carried out in the following manner: reject \( H_{r_s} \) if \( \theta_{0,r_s} \notin [w_{T,r_s} - c_1, \infty) \), for \( s = 1, \ldots, S \). Equivalently, reject \( H_{r_s} \) if \( w_{T,r_s} - \theta_{0,r_s} > c_1 \), for \( s = 1, \ldots, S \).

How should the value \( c_1 \) in the construction of the joint region (11) be chosen? Let \( \tilde{K} \) denote the index set that corresponds to the relabeled true hypotheses, that is,

\[s \in \tilde{K} \iff r_s \in I_0.\]

Ideally now, one would take \( c_1 = c_{\tilde{K}}(1 - \alpha, k, P) \), since this choice yields control of the \( k \)-FWE. To see why, assume w.l.o.g. that at least \( k \) hypotheses are true; otherwise, there is nothing to show. Then, with \( c_1 = c_{\tilde{K}}(1 - \alpha, k, P) \),

\[k\text{-FWE}_P = P\{\text{Reject at least } k \text{ of the } H_s : s \in I_0\} = P\{\text{Reject at least } k \text{ of the } H_{r_s} : s \in \tilde{K}\} = P\{\text{k-max}_{s \in \tilde{K}}(w_{T,r_s} - \theta_{r_s}) > c_{\tilde{K}}(1 - \alpha, k, P)\} \leq \alpha \text{ (by definition of } c_{\tilde{K}}(1 - \alpha, k, P)) \].

Unfortunately, the ideal choice \( c_1 = c_{\tilde{K}}(1 - \alpha, k, P) \) is not available for two reasons. First, the set \( \tilde{K} \) is unknown. Second, the probability mechanism \( P \) is unknown. The solution to the first problem is to replace \( \tilde{K} \) by \( \{1, \ldots, S\} \). Since \( \tilde{K} \subset \{1, \ldots, S\} \), it follows that \( c_{\tilde{K}}(1 - \alpha, k, P) \leq c_{\{1, \ldots, S\}}(1 - \alpha, k, P) \) and so the \( k \)-FWE is still controlled. The solution to the second problem is to replace \( P \) by an estimate \( \hat{P}_T \), that is, to apply the bootstrap. The choice of \( \hat{P}_T \) depends on context; see Appendix B of Romano and Wolf (2005c) for details. The cost of replacing \( P \) by \( \hat{P}_T \) is that control of the \( k \)-FWE is weakened to asymptotic control of the \( k \)-FWE. Combining the two solutions yields the choice \( \hat{c}_1 = c_{\{1, \ldots, S\}}(1 - \alpha, k, \hat{P}_T) \).

By continuing after the first step, more hypotheses can be rejected. Romano and Wolf (2005a) show that this increase in power does not come at the expense of sacrificing asymptotic control of the \( k \)-FWE. Denote by \( R_1 \) the number of rejections in the first step. If \( R_1 < k \), stop, since

\[\text{This region could also be called a generalized confidence region for reasons detailed below.}\]
it is ‘plausible’ that all rejected hypotheses are true ones. On the other hand, by controlling the $k$-FWE, if $R_1 \geq k$, we can be ‘confident’ that some of the rejected hypotheses are false. This ‘knowledge’ will now lead to smaller joint regions in subsequent steps, and hence to potentially further rejections, without sacrificing control of the $k$-FWE. So if $R_1 \geq k$, continue with the second step and construct a rectangular joint region for the vector $(\theta_{r_{R_1+1}}, \ldots, \theta_{r_S})'$ of the form

$$[w_{T, r_{R_1+1}} - c_2, \infty) \times \ldots \times [w_{T, r_S} - c_2, \infty).$$

(12)

Individual decisions are carried out analogously to before: reject $H_{r_s}$ if $\theta_{0, r_s} \notin [w_{T, r_s} - c_2, \infty)$, for $s = R_1 + 1, \ldots, S$.

How should the value $c_2$ in the joint region construction (12) be chosen? Again, the ideal choice $c_2 = c_K(1 - \alpha, k, P)$ is not available because $K$ and $P$ are unknown. Crucially, instead of replacing $K$ by $\{1, \ldots, S\}$, we can use information from the first step to arrive at a smaller value. Namely, if $P$ was known, this value would be given by

$$c_2 = \max\{c_K(1 - \alpha, k, P) : K = I \cup \{R_1 + 1, \ldots, S\}, I \subset \{1, \ldots, R_1\}, |I| = k - 1\}.$$

which is the maximum of a set of $\binom{R_1}{k-1}$ quantiles. The index set of any given quantile corresponds to all the hypotheses not rejected plus $k - 1$ out of the $R_1$ hypotheses that were rejected in the first step; and then one takes the largest such quantile for $c_2$. The intuition here is as follows. In order to ensure control of the $k$-FWE in the second step, $c_2$ must satisfy $c_2 \geq c_K(1 - \alpha, k, P)$. Assuming that $k$-FWE control was achieved in the first step, it is conceivable that up to $k - 1$ true hypotheses have been rejected so far. But, of course, we cannot know which of the rejected hypotheses might be true ones. So, to play it safe, one must look at all possible combinations of $k - 1$ rejected hypotheses, always together with the not rejected ones, and then take the largest of the resulting quantiles. Again, $P$ is unknown and so $c_2$ is not available in practice. Replacing $P$ by $\hat{P}_T$ yields the estimate

$$\hat{c}_2 = \max\{c_K(1 - \alpha, k, \hat{P}_T) : K = I \cup \{R_1 + 1, \ldots, S\}, I \subset \{1, \ldots, R_1\}, |I| = k - 1\}.$$

If no further hypotheses are rejected in the second step, stop. Otherwise, continue in this stepwise fashion until no more rejections occur. The following algorithm summarizes the procedure.

**Algorithm 4.1 (Basic $k$-StepM Method for One-Sided Setup)**

1. Relabel the strategies in descending order of the test statistics $w_{T, r}$: strategy $r_1$ corresponds to the largest test statistic and strategy $r_S$ to the smallest one.
2. For $1 \leq s \leq S$, if $\theta_{0,r_s} \notin [w_{T,r_s} - \hat{c}_1, \infty)$, reject the null hypothesis $H_{r_s}$. Here

$$\hat{c}_1 = c_{\{1, \ldots, S\}}(1 - \alpha, k, \hat{P}_T).$$

3. Denote by $R_1$ the number of hypotheses rejected. If $R_1 < k$, stop; otherwise let $j = 2$.

4. For $R_{j-1} + 1 \leq s \leq S$, if $\theta_{0,r_s} \notin [w_{T,r_s} - \hat{c}_j, \infty)$, reject the null hypothesis $H_{r_s}$. Here

$$\hat{c}_j = \max\{c_K(1 - \alpha, k, \hat{P}_T) : K = I \cup \{R_{j-1} + 1, \ldots, S\}, I \subset \{1, \ldots, R_{j-1}\}, |I| = k - 1\}. \quad (13)$$

5. (a) If no further hypotheses are rejected, stop.

(b) Otherwise, denote by $R_j$ the number of all hypotheses rejected so far and, afterwards, let $j = j + 1$. Then return to step 4.

The computation of the constants $\hat{c}_j$ via the bootstrap is detailed in Algorithm A.1 in the appendix. Let $J_T(P)$ denote the sampling distribution under $P$ of $\sqrt{T}(W_T - \theta)$; and let $J_T(\hat{P}_T)$ denote the sampling distribution under $\hat{P}_T$ of $\sqrt{T}(W_T^* - \hat{\theta}_T)$. Here, $\hat{\theta}_T$ is an estimate of $\theta$ based on $\hat{P}_T$.\(^9\) Romano and Wolf (2005a) show that a sufficient condition for the basic $k$-StepM method to control the $k$-FWE asymptotically is the following.

**Assumption 4.1** Let $P$ denote the true probability mechanism and let $\hat{P}_T$ denote an estimate of $P$ based on the data $X_T$. Assume that $J_T(P)$ converges in distribution to a limit distribution $J(P)$, which is continuous. Further assume that $J_T(\hat{P}_T)$ consistently estimates this limit distribution: $\rho(J_T(\hat{P}_T), J(P)) \to 0$ in probability for any metric $\rho$ metrizing weak convergence.

We now describe how the basic StepM method is modified for the two-sided setup (2). The crux is that the multivariate rectangular joint regions are now the Cartesian products of two-sided intervals rather than one-sided intervals.

To this end, for any $K \subset \{1, \ldots, S\}$, define

$$c_{K,1\mid}(1 - \alpha, k, P) = \inf\{x : P\{k-\text{max}_{s \in K}|w_{T,r_s} - \theta_{r_s}| \leq x\} \geq 1 - \alpha\}. \quad (14)$$

That is, $c_{K,1\mid}(1 - \alpha, k, P)$ is the smallest $1 - \alpha$ quantile of the 'two-sided' sampling distribution under $P$ of $k-\text{max}_{s \in K}|w_{T,r_s} - \theta_{r_s}|$.

The following algorithm summarizes our stepwise method modified for the two-sided setup.

\(^9\)Usually, one can take $\hat{\theta}_T = \theta(\hat{P}_T)$.
Algorithm 4.2 (Basic k-StepM Method for Two-Sided Setup)

1. Relabel the strategies in descending order of the absolute test statistics $|w_{T,s}|$: strategy $r_1$ corresponds to the largest absolute test statistic and strategy $r_S$ to the smallest one.

2. For $1 \leq s \leq S$, if $\theta_{0,r_s} \not\in [w_{T,r_s} \pm \hat{c}_{1,|s|}]$, reject the null hypothesis $H_{r_s}$. Here
\[
\hat{c}_{1,|s|} = c_{1,\ldots,S,|s|}(1 - \alpha, k, \hat{P}_T).
\]

3. Denote by $R_1$ the number of hypotheses rejected. If $R_1 < k$, stop; otherwise let $j = 2$.

4. For $R_{j-1} + 1 \leq s \leq S$, if $\theta_{0,r_s} \not\in [w_{T,r_s} \pm \hat{c}_{j,|s|}]$, reject the null hypothesis $H_{r_s}$. Here
\[
\hat{c}_{j,|s|} = \max\{c_{K,|s|}(1 - \alpha, k, \hat{P}_T) : K = I \cup \{R_{j-1} + 1, \ldots, S\}, I \subset \{1, \ldots, R_{j-1}\}, |I| = k - 1\}.
\]

5. (a) If no further hypotheses are rejected, stop.

(b) Otherwise, denote by $R_j$ the number of all hypotheses rejected so far and, afterwards, let $j = j + 1$. Then return to step 4.

The computation of the constants $\hat{c}_{j,|s|}$ via the bootstrap is detailed in Algorithm A.2 in the appendix. A sufficient condition for the basic k-StepM method to asymptotically control the $k$-FWE in the two-sided setup is also given by Assumption 4.1.

Remark 4.1 The procedure we suggest is based on symmetric two-sided confidence intervals. Alternatively, it could be based on equal-tailed two-sided confidence intervals. See Hall (1992) for a comparison of these interval types in the context of inference for univariate parameters.

4.3.2 Studentized Method

We now describe how to modify the $k$-StepM method when studentized test statistics are used instead. Ample motivation for the desirability of studentization in the context of FWE control is provided by Hansen (2005) and Romano and Wolf (2005c). Their reasons carry over to $k$-FWE control.

Again, we start with the one-sided setup (1). Analogously to (10), define
\[
d_K(1 - \alpha, k, P) = \inf\{x : P\{k\text{-max}(|w_{T,r_s} - \theta_{r_s}|/\hat{\sigma}_{T,r_s} : s \in K) \leq x\} \geq 1 - \alpha\}.
\]

15
Our stepwise method is then summarized by the following algorithm.

Algorithm 4.3 (Studentized k-StepM Method for One-Sided Setup)

1. Relabel the strategies in descending order of the studentized test statistics $z_{T,s}$: strategy $r_1$ corresponds to the largest test statistic and strategy $r_S$ to the smallest one.

2. For $1 \leq s \leq S$, if $\theta_{0,r_s} \notin [w_{T,r_s} - \hat{\sigma}_{T,r_s} \hat{\Delta}_1, \infty)$, reject the null hypothesis $H_{r_s}$. Here

$$\hat{\Delta}_1 = d_{\{1,\ldots,S\}}(1 - \alpha, k, \hat{P}_T).$$

3. Denote by $R_1$ the number of hypotheses rejected. If $R_1 < k$, stop; otherwise let $j = 2$.

4. For $R_{j-1} + 1 \leq s \leq S$, if $\theta_{0,r_s} \notin [w_{T,r_s} - \hat{\sigma}_{T,r_s} \hat{\Delta}_j, \infty)$, reject the null hypothesis $H_{r_s}$. Here

$$\hat{\Delta}_j = \max\{d_K(1 - \alpha, k, \hat{P}_T) : K = \{1\} \cup \{R_{j-1} + 1, \ldots, S\}, I \subset \{1, \ldots, R_{j-1}\}, |I| = k - 1\}. \quad (17)$$

5. (a) If no further hypotheses are rejected, stop.

(b) Otherwise, denote by $R_j$ the number of all hypotheses rejected so far and, afterwards, let $j = j + 1$. Then return to step 4.

The computation of the constants $\hat{\Delta}_j$ via the bootstrap is detailed in Algorithm A.3 in the appendix.

The modification to the two-sided setup (2) is now quite obvious. Analogously to (16), define

$$d_{K,\{\cdot\}}(1 - \alpha, k, P) = \inf\{x : P\{k-\max(|w_{T,r_s} - \theta_{r_s}|/\hat{\sigma}_{T,r_s} : s \in K) \leq x\} \geq 1 - \alpha\}. \quad (18)$$

The algorithm can then be summarized as follows.

Algorithm 4.4 (Studentized k-StepM Method for Two-Sided Setup)

1. Relabel the strategies in descending order of the absolute studentized test statistics $|z_{T,s}|$: strategy $r_1$ corresponds to the largest absolute studentized test statistic and strategy $r_S$ to the smallest one.

2. For $1 \leq s \leq S$, if $\theta_{0,r_s} \notin [w_{T,r_s} \pm \hat{\sigma}_{T,r_s} \hat{\Delta}_{1,\{\cdot\}}, \infty)$, reject the null hypothesis $H_{r_s}$. Here

$$\hat{\Delta}_{1,\{\cdot\}} = d_{\{1,\ldots,S\},\{\cdot\}}(1 - \alpha, k, \hat{P}_T).$$
3. Denote by $R_1$ the number of hypotheses rejected. If $R_1 < k$, stop; otherwise let $j = 2$.

4. For $R_{j-1} + 1 \leq s \leq S$, if $\theta_{0,r_s} \notin [w_{T,r_s} \pm \hat{\sigma}_{T,r_s} \hat{d}_{j,|I|}]$, reject the null hypothesis $H_{r_s}$. Here

$$
\hat{d}_{j,|I|} = \max\{d_{K,|I|}(1 - \alpha, k, \hat{P}_T) : K = I \cup \{R_{j-1} + 1, \ldots, S\}, I \subset \{1, \ldots, R_{j-1}\}, |I| = k - 1\}.
$$

(19)

5. (a) If no further hypotheses are rejected, stop.

(b) Otherwise, denote by $R_j$ the number of all hypotheses rejected so far and, afterwards, let $j = j + 1$. Then return to step 4.

The computation of the constants $\hat{d}_{j,|I|}$ via the bootstrap is detailed in Algorithm A.4 in the appendix.

A slightly stronger version of Assumption 4.1 is needed to prove the validity of the studentized method. Again, let $X^*_T$ denote a data matrix generated from probability mechanism $\hat{P}_T$. The basic test statistics computed from $X^*_T$ are denoted by $w^*_{T,s}$. Their corresponding standard errors, also computed from $X^*_T$, are denoted by $\hat{\sigma}_{T,s}^*$. Romano and Wolf (2005a) do not explicitly discuss the case of studentized statistics. However, it is straightforward to show that a sufficient condition for the studentized $k$-StepM method to control the $k$-FWE asymptotically, both in the one-sided and the two-sided setup, is the following.

**Assumption 4.2** In addition to Assumption 4.1, assume the following condition. For each $1 \leq s \leq S$, both $\sqrt{T} \hat{\sigma}_{T,s}$ and $\sqrt{T} \hat{\sigma}_{T,s}^*$ converge to a (common) positive constant $\sigma_s$ in probability.

**Remark 4.2 (Operative Method)** The computation of the constants $\hat{c}_j$, $\hat{d}_{j,|I|}$, $\hat{d}_j$, and $\hat{\sigma}_{T,s}^*$ in (13), (15), (17), and (19), respectively, may be very expensive in case $\binom{R_j - 1}{k-1}$ is large. In such cases, we suggest the following shortcut. Pick a user-defined number $N_{max}$, say $N_{max} = 50$ and let $N^*$ be the largest integer for which $\binom{N^*}{k-1} \leq N_{max}$. The constant $\hat{c}_j$, say, is then computed as

$$
\hat{c}_j = \max\{\hat{c}_K(1 - \alpha, k, \hat{P}_T) : K = I \cup \{R_{j-1} + 1, \ldots, S\}, I \subset \{R_j - N^* + 1, \ldots, R_j\}, |I| = k - 1\},
$$

and similarly for the constants $\hat{c}_{j,|I|}$, $\hat{d}_j$, and $\hat{d}_{j,|I|}$. That is, we maximize over subsets $I$ not necessarily of the entire index set of previously rejected hypotheses but only of the index set corresponding to the $N^*$ least significant hypotheses rejected so far. Note that this shortcut does not affect the asymptotic control of the $k$-FWE even if $N_{max} = 1$ is chosen, resulting in $N^* = k - 1$ and

$$
\hat{c}_j = \hat{c}_{\{R_j - k, \ldots, S\}}(1 - \alpha, k, \hat{P}_T).
$$
Nevertheless, in the interest of better $k$-FWE control in finite samples, we suggest to choose $N_{\text{max}}$ as large as possible.

**Remark 4.3** All methods presented in this section can be modified in the following sense while still preserving (asymptotic) control of the $k$-FWE: reject the $k - 1$ ‘most significant’ hypotheses no matter what. This means sort the hypotheses in the order of either ascending $p$-values or descending test statistics to get $H_{r_1}, \ldots, H_{r_5}$; then reject $H_{r_1}, \ldots, H_{r_{k-1}}$ irrespective of the data. Let $R$ denote the number of rejections made by the multiple testing method (before modification). If $R < k$, then the modified method will reject $k - 1$ hypotheses, that is, a potentially greater number. If $R \geq k$, then the modified method will reject $R$ hypotheses, that is, the same number. However, it is counterintuitive to reject hypotheses irrespective of the data and certainly we would also impose the minimal requirement to not reject any hypothesis when the corresponding marginal $p$-value exceeds $\alpha$.

**4.3.3 The StepM Method**

Naturally, the StepM Method of Romano and Wolf (2005c) can be considered a special case of the $k$-StepM method by choosing $k = 1$. However, it should be pointed out that the computations are much simplified compared to the case $k > 1$. The reason is that if for the StepM method some hypotheses are rejected in the first step, then for the computation of the values $\hat{c}_j$ and $\hat{d}_j$, $j = 2, 3, \ldots$, one may assume that all hypotheses rejected so far are false ones.\(^{10}\) As a result, in the $j$th step, one does not have to compute the maximum of a set of $\binom{R_{j-1}}{k-1}$ estimated quantiles but rather only a single estimated quantile.

The following algorithm is the simplified Version of Algorithm 4.1 for the special case $k = 1$. The simplified versions of Algorithms 4.2–4.4 are analogous.

**Algorithm 4.5 (Basic StepM Method for One-Sided Setup)**

1. Relabel the strategies in descending order of the test statistics $w_{T,s}$; strategy $r_1$ corresponds to the largest test statistic and strategy $r_S$ to the smallest one.

2. For $1 \leq s \leq S$, if $\theta_{0,s} \notin [w_{T,s} - \hat{c}_1, \infty)$, reject the null hypothesis $H_{r_s}$. Here

\[
\hat{c}_1 = c_{(1, \ldots, S)}(1 - \alpha, 1, \hat{P}_T).
\]

\(^{10}\)If a true hypothesis has been rejected so far, then the FWE criterion has already been violated and, therefore, the rejection of further true hypotheses will not do any additional harm.
3. Denote by $R_1$ the number of hypotheses rejected. If $R_1 = 0$, stop; otherwise let $j = 2$.

4. For $R_{j-1} + 1 \leq s \leq S$, if $\theta_{0,r_s} \not\in [w_{T,r_s} - \hat{c}_j, \infty)$, reject the null hypothesis $H_{r_s}$. Here

\[ \hat{c}_j = c_{\{R_{j-1} + 1, \ldots, s\}}(1 - \alpha, 1, \hat{p}_T). \]

5. (a) If no further hypotheses are rejected, stop.

(b) Otherwise, denote by $R_j$ the number of all hypotheses rejected so far and, afterwards, let $j = j + 1$. Then return to step 4.

5 Methods Controlling the FDR

In many applications, one might be willing to tolerate a larger number of false rejections if there are a larger number of total rejections. In other words, one might be willing to tolerate a certain (small) proportion of false rejections out of the total rejections. This suggests to base error control on the false discovery proportion (FDP). Let $F$ be the number of false rejections made by a multiple testing method and let $R$ be the total number of rejections. Then the FDP is defined as follows:

\[ \text{FDP} = \begin{cases} \frac{F}{R} & \text{if } R > 0 \\ 0 & \text{if } R = 0 \end{cases}. \]

Benjamini and Hochberg (1995) propose to control $E_P(\text{FDP})$, the expected value under $P$ of the FDP, which they coined the false discovery rate (FDR). A multiple testing method is said to control the FDR at level $\gamma$ if $\text{FDR}_P = E_P(\text{FDP}) \leq \gamma$ for any sample size $T$ and for any $P$. A multiple testing method is said to control the FDR asymptotically at level $\gamma$ if $\limsup_{T \to \infty} \text{FDR}_P \leq \gamma$ for any $P$. Methods that control the FDR in finite samples can typically only be derived in special circumstances.

The stepwise method of Benjamini and Hochberg (1995) is based on individual $p$-values. The $p$-values are ordered from smallest to largest, $\hat{p}_{T,(1)} \leq \hat{p}_{T,(2)} \leq \ldots \leq \hat{p}_{T,(S)}$, with their corresponding null hypotheses labeled accordingly, $H_{(1)}$, $H_{(2)}$, $\ldots$, $H_{(S)}$. Then define

\[ j^* = \max \{ j : \hat{p}_{T,(j)} \leq \gamma_j \} \quad \text{where} \quad \gamma_j = \frac{j}{S} \gamma \quad (20) \]

and reject $H_{(1)}, \ldots, H_{(j^*)}$. If no such $j$ exists, reject no hypotheses. This is an example of a stepup method. It starts with examining the least significant hypothesis, $H_{(S)}$, and then moves ‘up’ to the more significant hypotheses if $\hat{p}_{T,(S)} > \gamma$. 

19
Benjamini and Hochberg (1995) prove that their method controls the FDR if condition (8) holds and, in addition, the \( p \)-values are independent of each other. Benjamini and Yekutieli (2001) show that independence can be replaced by a more general 'positive regression dependency'; see their paper for the exact definition. As a result, it can be proven that, under the dependence condition of Benjamini and Yekutieli (2001), the method of Benjamini and Hochberg (1995) asymptotically controls the FDR if condition (9) hold. On the other hand, (asymptotic) control of the Benjamini and Hochberg (1995) method under an arbitrary dependence structure of the \( p \)-values cannot be demonstrated. Benjamini and Yekutieli (2001) show that this more general control can be achieved if the constants \( \gamma_j \) in (20) are replaced by

\[
\gamma_j = \frac{j}{S} \frac{\gamma}{C_S} \quad \text{where} \quad C_k = \sum_{s=1}^{k} \frac{1}{s^2}.
\]

Note that \( C_S \approx \log(S) + 0.5 \) and so this method can have much less power than the original Benjamini and Hochberg (1995) method. For example, when \( S = 1,000 \), then \( C_S = 7.49 \).

Leaving aside the issue of sufficient conditions to ensure control of the FDR, there is a shortcoming with this particular generalized error rate that is often ignored. The FDR is the mean of the FDP, that is, a central tendency of the sampling distribution of the FDP. Therefore, even if the FDR is controlled at level \( \gamma \), in a given application, the realized FDP could be quite far away from \( \gamma \). How likely this is depends on the sampling variability of the FDP which is unknown in practice.\(^\text{11}\) Korn et al. (2004) provide some simulations which highlight this shortcoming; also see Section 8.

### 6 Methods Controlling the FDP

Often, one would like to be able to make a statement concerning the realized FDP in a given application. Concretely, one would like to control the FDP in the sense that \( P\{\text{FDP} > \gamma \} \) where \( \gamma \in [0,1] \) is a user-defined number. Typical values are \( \gamma = 0.05 \) and \( \gamma = 0.1 \); the choice \( \gamma = 0 \) corresponds to control of the FWE.

A multiple testing method is said to control the FDP at level \( \alpha \) if \( P\{\text{FDP} > \gamma \} \leq \alpha \) for any sample size \( T \) and for any \( P \). A multiple testing method is said to control the FDP asymptotically at level \( \alpha \) if \( \limsup_{T \to \infty} P\{\text{FDP} > \gamma \} \leq \alpha \) for any \( P \). Methods that control the FDP in finite samples can typically only be derived in special circumstances.

As an alternative to controlling the FDR, which is the expected value of the FDP, we propose

\(^{11}\text{Obviously, some very crude bounds could be obtained using Markov's inequality or variants thereof.}\)
controlling the median of the FDP. Obviously, if we achieve \( P\{\text{FDP} > \gamma\} \leq 0.5 \), then the median FDP is bounded above by \( \gamma \). So choosing \( \alpha = 0.5 \) for the methods in this section, asymptotically controls the median FDP under an arbitrary dependence structure of the \( p \)-values (or test statistics).

We now describe how some of the methods of Section 3 can be generalized to achieve (asymptotic) control of the FDP. Of course, since our goal is to reject as many false hypotheses as possible, in the end we shall recommend the generalization of the StepM method.

### 6.1 Generalization of the Holm Method

Lehmann and Romano (2005) develop a stepdown method based on individual \( p \)-values. The \( p \)-values are ordered from smallest to largest, \( \hat{p}_{T,(1)} \leq \hat{p}_{T,(2)} \leq \ldots \leq \hat{p}_{T,(S)} \), with their corresponding null hypotheses labeled accordingly, \( H_{(1)}, H_{(2)}, \ldots, H_{(S)} \). Then \( H_{(s)} \) is rejected at level \( \alpha \) if \( \hat{p}_{T,(j)} \leq \alpha_j \) for \( j = 1, \ldots, s \), where

\[
\alpha_j = \frac{[\gamma j] + 1)\alpha}{S + [\gamma j] + 1 - j}.
\]

Here, for a real number \( x \), \( \lfloor x \rfloor \) denotes the greatest integer which is smaller than or equal to \( x \).

It can be proven that this method provides asymptotic control of the FDP if condition (9) holds. Moreover, this method provides finite-sample control of the FDP if condition (8) holds and the \( p \)-values are independent, or at least positively dependent in a certain sense; see Lehmann and Romano (2005). Lehmann and Romano (2005) also show that if one modifies this method by replacing \( \alpha_j \) by

\[
\alpha'_j = \frac{\alpha_j}{C_{[\gamma S]+1}} \quad \text{where} \quad C_k = \sum_{s=1}^{k} \frac{1}{s},
\]

then the resulting stepdown procedure controls the FDP under no dependence assumptions on the \( p \)-values. This method has since been improved by Romano and Shaikh (2005) in that the constant \( C_{[\gamma S]+1} \) has been replaced by a smaller one, while still maintaining finite-sample control under assumption (8) and asymptotic control under assumption (9). A similar stepup procedure is derived in Romano and Shaikh (2006).

### 6.2 Generalization of the StepM Method

The crux of our procedure is to sequentially apply the \( k \)-StepM method, \( k = 1, 2, 3, \ldots \), until a stopping rule indicates termination. The appropriate variant of the \( k \)-StepM method is dictated by the nature of the multiple testing problem, one-sided versus two-sided, and the choice of test
statistics, basic versus studentized. For example, the one-sided setup (1) in combination with studentized test statistics calls for Algorithm 4.3.

To develop the idea, consider controlling $P\{\text{FDP} > 0.1\}$. We start out by applying the 1-StepM method, that is, by controlling the FWE. Denote by $N_1$ the number of hypotheses rejected. Due to the FWE control one can be confident that no false rejection has occurred and that, in return, the FDP has been controlled. Consider now rejecting $H_{(N_1+1)}$, the next ‘most significant’ hypothesis. Of course, if $H_{(N_1+1)}$ is false, there is nothing to worry about, so suppose $H_{(N_1+1)}$ is true. Assuming FWE control in the first step, the FDP upon rejection of $H_{(N_1+1)}$ then becomes $1/(N_1 + 1)$, which is greater than 0.1 if and only if $N_1 < 9$. So if $N_1 \geq 9$ we can reject one true hypothesis and still avoid FDP > 0.1. This suggests to stop if $N_1 < 9$ and to otherwise apply the 2-StepM method which, by design, should not reject more than one true hypothesis. Denote the total number of hypotheses rejected by the 2-StepM method by $N_2$. Reasoning similarly to before, if $N_2 < 19$, we stop and otherwise we apply the 3-StepM method. This procedure is continued until $N_j < 10j - 1$ at some point.

The following algorithm summarizes the method for arbitrary $\gamma$.

**Algorithm 6.1 (FDP-StepM Method)**

1. Let $j = 1$ and let $k_1 = 1$.

2. Apply the $k_j$-StepM method and denote by $N_j$ the number of hypotheses rejected.

3. (a) If $N_j < k_j/\gamma - 1$, stop.

   (b) Otherwise, let $j = j + 1$ and, afterwards, let $k_j = k_{j-1} + 1$. Then return to step 2.

Romano and Wolf (2005a) show that a sufficient condition for the FDP-StepM method to provide asymptotic control of the FDP is Assumption 4.1 in case the underlying $k$-StepM method uses basic test statistics. Similarly, it can be proven that a sufficient condition for the FDP-StepM method to provide asymptotic control of the FDP is Assumption 4.2 in case the underlying $k$-StepM method uses studentized test statistics.

7 Applications to Model Selection

This section briefly discusses how control of generalized error rates can apply to the problem of model selection. Unfortunately, the term ‘model selection’ is rather vague and can mean different things depending on context. Therefore, we consider various notions.
White (2000) studies the problem of comparing a large number of (forecast) models to a common (forecast) benchmark. In this context, model selection is the challenge of deciding which models are superior to the benchmark. Therefore, in this context, model selection becomes a special case of Example 2.1 by interpreting (forecast) models as strategies. White (2000) proposes control of the FWE, but when the number of strategies is very large this criterion can be too strict and a generalized error rate may be more appropriate. Some empirical applications based on the FWE when the number of strategies is very large are the following. Sullivan et al. (1999), White (2000), and Sullivan et al. (2001) all fail to find any outperforming strategies when comparing a large number, $S$, of trading strategies against the benchmark of ‘buy and hold’. The numbers of trading strategies considered are $S = 7,846$, $S = 3,654$, and $S = 9,452$, respectively. Hansen (2005) fails to find any outperforming strategies when comparing $S = 3,304$ strategies to forecast inflation against the benchmark of ‘last period’s inflation’. On the other hand, when he restricts attention to a smaller universe of $S = 352$ strategies, some outperformers are detected. It appears that when the number of strategies is in the thousands, controlling the FWE becomes too stringent.

The task of constructing an ‘optimal’ forecast provides another notion. Imagine several forecasting strategies are available to forecast a quantity of interest. As described in Timmermann (2006, Chapter 6): (i) choosing the lone strategy with the best track record is often a bad idea; (ii) simple forecasting schemes, such as equal-weighting various strategies, are hard to beat; and (iii) trimming off the worst strategies is often required. In this context, model selection is the challenge of identifying the ‘worst strategies’. A sensible approach is the following. First, one labels those strategies as the ‘worst strategies’ which underperform a suitable benchmark.\footnote{For example, when forecasting inflation a suitable, simple-minded benchmark might be last period’s inflation.} Second, one is now back again in Example 2.1 except that the individual parameters need to be defined in such a way that $\theta_s \leq 0$ if and only if the $s$th strategy does not underperform the benchmark. Typically, this can be achieved by defining the parameters according to Example 2.1 and then reversing their sign.

In many empirical applications, a large-dimensional regression model is estimated and the question becomes which explanatory variables are the ‘important’ ones. In this context, model selection is the challenge of identifying the non-zero regression coefficients; see Example 2.2. The unfortunate common practice is identification based on individual $p$-values, ignoring the multiple testing problem altogether.\footnote{For example, it is common to provide tables where the ‘important’ explanatory variables are identified via superscript stars: one star if significant at level 10%, two stars if significant at level 5%, and three stars if significant at level 1%. (Where the levels are for individual tests always.)} As a result, one typically identifies too many variables as ‘important’. For example, if there are 100 variables under test, all of which are ‘unimportant’, then, based on comparing individual $p$-values to the level $\alpha = 5\%$, one would expect to falsely identify five variables
as 'important'. On the other hand, dealing with the multitude of test by applying the FWE can be too strict, especially when the number of explanatory variables is very large. As a result, one may easily overlook 'important' variables. The sensible solution is therefore to employ a suitable generalized error rate, such as controlling the (median) FDP. Note that the estimated regression coefficients may dependent on each other in a way that violates the positive regression dependency assumption, and so the validity of the FDR procedure of Benjamini and Hochberg (1995) is not guaranteed.

Related to the model selection notion of the previous paragraph, though more complex, is the problem of determining which explanatory variables to keep in a final model, say for prediction purposes. This problem is commonly known as 'subset selection' and many popular techniques exist, such as pretesting methods, stepwise selection (forward or backward), the application of information criteria such as AIC or BIC, and principle components regression. See Draper and Smith (1998) or Hastie et al. (2001) for details. An explicit use of tests of multivariate parameters as a means of consistent variable selection can be found in Pötscher (1983) and Bauer et al. (1988). Another popular technique for 'subset selection' is general-to-specific modeling; see Campos et al. (2005) for an introduction. As a part of the procedure, individual variables are kept in the model based on so-called simplification tests where individual p-values are compared to a (common) significance level \( \alpha \). The choice of this level appears as much of an art as a science. For example, Krolzig and Hendry (2001) discuss how it is advantageous to choose a small level \( \alpha \) when there are many irrelevant explanatory variables. However, they do not address the question of how one is to know whether this is the case in practice. The 'optimal' level \( \alpha \) for the individual tests depends not only on the number of explanatory variables, which is known, but also on the number of them that are irrelevant and the dependence structure of the regression coefficient estimates, both of which are unknown. Therefore, a viable alternative may be to consider the simplification tests as a multiple testing problem in conjunction with a generalized error rate such as controlling the (median) FDP. Such an approach can implicitly account both for the number of irrelevant variables and the dependence structure of the regression coefficient estimates.

Jensen and Cohen (2000) discuss multiple comparisons in induction algorithms. In this context, model selection is the challenge of deciding which variables to include in an AI (artificial intelligence) model for prediction and classification purposes. They describe how a procedure ignoring the multiple testing problem leads to undesirable effects such as overfitting, that is, the inclusion of too many variables in the model. Control of a generalized error rate may therefore be desirable. Moreover, Jensen and Cohen (2000) show in some simulations that multiple testing procedures that do not account for the dependence structure of the test statistics, such as Bonferroni, can work well when the dependence structure is absent or weak but work poorly when the dependence structure is
noticeable. Hence, it is desirable to employ a procedure that accounts for the dependence structure.

Abramovich and Benjamini (1996) and Abramovich et al. (2005) study the problem of recovering an \( S \)-dimensional vector observed in white noise, where \( S \) is large and the vector is known to be sparse. Abramovich et al. (2005) discuss various definitions of sparseness, the most intuitive one being the proportion of the non-zero entries of the vector. In this context, model selection is the challenge of deciding which entries are non-zero in order to 'optimally' estimate the vector.\(^{14}\) The suggested solution is to consider the problem as a suitable multiple testing problem where the individual hypotheses test whether the entries of the vector are zero or non-zero (and so the hypotheses are two-sided). Then the FDR criterion is employed to account for the multitude of tests. Abramovich et al. (2005) show that this approach based on the FDR enjoys optimality properties, but their asymptotic framework is somewhat restrictive.\(^{15}\) In addition, the error terms are assumed independent of each other. As an alternative, which is also robust against dependent errors, one might control the (median) FDP instead.

Recently, Buena et al. (2006) show how testing using FDR control can be used to produce consistent variable selection even in high dimensional models. Of course, other measures of error control can similarly be exploited.

We also mention a notion of model selection that does not fit in our framework. Again, imagine several forecasting strategies are available to forecast a quantity of interest. But now the question is which of those strategies is the 'best'. In this context, model selection is the challenge of identifying the 'best' model out of a universe of 'candidate' models. Needless to say, given a finite amount of data, the 'best' model cannot determined with certainty. Therefore, the suggested solution consists of constructing a model confidence set, that is, a data-dependent collection of models which will contain the 'best' model with a pre-specified probability, at least asymptotically. For related work see Shimodaira (1998), Hansen et al. (2003), Hansen et al. (2005), and the references therein.

While the above discussion reveals that multiple hypothesis testing methods may be useful in the model building process, the problem of inference for parameters of a data-based model is crucial and challenging. For recent entries to the literature on inference after model selection, see Shen et al. (2004) and Kabaila and Leeb (2006) as well as the references in these works.

\(^{14}\)Here optimality is defined in an asymptotic minimax sense; see Abramovich et al. (2005) for details.

\(^{15}\)For instance, they assume that the sparsity tends to zero, so the limiting model for the vector is the one of a 'black object' (where all entries are equal to zero).
8 Simulation Study

This section presents a small simulation study in the context of testing population means. We generate independent random vectors $X_1, \ldots, X_T$ from an $s$-dimensional multivariate normal distribution with mean vector $\theta = (\theta_1, \ldots, \theta_S)'$, where $T = 100$ and $S = 500$. The null hypotheses are $H_s : \theta_s \leq 0$ and the alternative hypotheses are $H_s : \theta_s > 0$, so we are in the one-sided setup (1). The studentized test statistics are $z_{T,s} = w_{T,s}/\hat{\sigma}_{T,s}$, where

$$w_{T,s} = \bar{X}_{., s} = \frac{1}{T} \sum_{t=1}^{T} X_{t,s} \quad \text{and} \quad \hat{\sigma}_{T,s}^2 = \frac{1}{T(T-1)} \sum_{t=1}^{T} (X_{t,s} - \bar{X}_{., s})^2 .$$

The individual means $\theta_s$ are equal to either 0 or 0.25. The number of means equal to 0.25 is 0, 100, 200, or 400. The covariance matrix is of the common correlation structure: $\sigma_{s,s} = 1$ and $\sigma_{s,j} = \rho$ for $s \neq j$. We consider the values $\rho = 0$ and $\rho = 0.5$. Other specifications of the covariance matrix do not lead to results that are qualitatively different; see Romano and Wolf (2005a).

We include the following multiple testing procedures in the study. The value of $k$ is $k = 10$. The nominal level is $\alpha = 0.05$, unless indicated otherwise.


- **(k-gH)** The $k$-FWE generalized Holm procedure described in Subsection 4.2, where the individual $p$-values are derived from $z_{T,s} \sim t_{T-1}$ under $\theta_s = 0$.

- **(k-StepM)** The studentized $k$-StepM construction described in Subsection 4.3. This procedure is based on the operative method with $N_{\text{max}} = 50$, see Remark 4.2.

- **(FDP-LR$_{0.1}$)** The FDP procedure of Lehmann and Romano (2005) with $\gamma = 0.1$ described in Subsection 6.1.

- **(FDP-StepM$_{0.1}$)** The studentized FDP-StepM construction described in Subsection 6.2 with $\gamma = 0.1$.

- **(FDP-StepM$_{0.1}^{\text{med}}$)** The studentized FDP-StepM construction described in Subsection 6.2 with $\gamma = 0.1$ but nominal level $\alpha = 0.5$. Therefore, this procedure asymptotically controls the median FDP to be bounded above by $\gamma = 0.1$.

- **(FDR-BH$_{0.1}$)** The FDR construction of Benjamini and Hochberg (1995) described in Section 5 with $\gamma = 0.1$.
The performance criteria are (i) the empirical $k$-FWEs and FDPs, compared to the nominal level $\alpha = 0.05$ (or $\alpha = 0.5$ for the method controlling the median FDP), and the empirical FDRs; and (ii) the average number of false hypotheses rejected. The results are presented in Table 1. They can be summarized as follows.

- All methods provide satisfactory finite sample control of their respective $k$-FWE, FDP, or FDR criteria.

- By controlling a generalized error rate, the power is often much improved compared to FWE control.

- The methods that implicitly account for the dependence structure of the test statistics are more powerful than the ‘worst case’ methods based on individual p-values: 10-StepM is more powerful than 10-gH and FDP-StepM$_{0.1}$ is more powerful than FDP-LR$_{0.1}$.

- The methods controlling a central tendency of the FDP are more powerful than the methods controlling $P\{\text{FDP} > 0.1\}$: FDP-StepM$_{0.1}^{\text{Med}}$ and FDR$_{0.1}$ are more powerful than FDP-LR$_{0.1}$ and FDP-StepM$_{0.1}$.

Given the last observation, one may be tempted to prefer FDP-StepM$_{0.1}^{\text{Med}}$ over FDP-StepM$_{0.1}$ always. However, it should be understood that these two approaches are philosophically different. If $P\{\text{FDP} > 0.1\} \leq 0.05$ is achieved, then, in a given application, one can be 95% confident that the realized FDP is at most 0.1. On the other hand, if $P\{\text{FDP} > 0.1\} \leq 0.5$ is achieved (i.e., control of the median FDP), then, in a given application, one can only be 50% confident that the realized FDP is at most 0.1. So, there may be a good chance that the realized FDP is greater than 0.1, and perhaps by quite a bit. The problem is similar with FDR control because it also controls only a central tendency of the FDP, namely the mean.

To examine this issue, we look at the sampling distribution of the FDP when the median FDP and the FDR are controlled. Figure 1 summarizes the distribution of the realized FDPs for various scenarios via boxplots. It can be seen that, while median FDP control and FDR control are achieved, the variation of the sampling distributions is considerable, especially for the case of common correlation $\rho = 0.5$. As a result, the realized FDP may well be quite above $\gamma = 0.1$.  

27
9 Empirical Applications

9.1 Hedge Fund Evaluation

The data set we consider is similar to one in Romano and Wolf (2005c). The difference is that we focus on a shorter time horizon, thereby increasing the number of funds under study. Our universe consists of all hedge funds in the Center for International Securities and Derivatives Markets (CISDM) database that have a complete return history from 01/1994 until 12/2003.

All returns are net of management and incentive fees, so they are the returns obtained by the investors. As in Romano and Wolf (2005c), we benchmark the funds against the risk-free rate\(^{16}\), and all returns are log returns. So we are in the situation of Example 2.1.(a). It is well known that hedge fund returns, unlike mutual fund returns, tend to exhibit non-negligible serial correlations; see, for example, Lo (2002) and Kat (2003). Accordingly, one has to account for this time series nature in order to obtain valid inference. Studentization for the original data uses a kernel variance estimator based on the prewhitened QS kernel and the corresponding automatic choice of bandwidth of Andrews and Monahan (1992). The bootstrap method is the circular block bootstrap, based on \( M = 5,000 \) repetitions. The studentization in the bootstrap world uses the corresponding `natural' variance estimator; for details, see Götze and Künsch (1996) or Romano and Wolf (2003). The block sizes for the circular bootstrap are chosen via Algorithm A.5. The semi-parametric model \( \hat{P}_T \) used in this algorithm is a VAR(1) model in conjunction with bootstrapping the residuals.\(^{17}\)

There are 210 funds in the CISDM database with a complete return history from 01/1994 until 12/2003, and the number of monthly observations is \( T = 120 \). However, one fund is deleted from the universe due to a highly unusual return distribution\(^{18}\), so the number of funds included in the study is \( S = 209 \) in the end. Table 2 lists the ten largest basic and studentized test statistics, together with the corresponding hedge funds. Similar to the analysis of Romano and Wolf (2005c), the two lists are almost completely disjoint; only the fund JMG Capital Partners appears in both lists.

We now use the various multiple testing methods to identify hedge funds that outperform the risk-free rate, starting with the the Holm procedure and its generalizations as well as the FDR procedure of Benjamini and Hochberg (1995), all of which are based on individual \( p \)-values

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\(^{16}\)The risk-free rate is a simple and widely accepted benchmark. But, of course, our methods also apply to alternative benchmarks such as hedge fund indices or multi-factor hedge fund benchmarks; for example, see Kosowski et al. (2005).

\(^{17}\)To account for leftover dependence not captured by the VAR(1) model, we use the stationary bootstrap with average block size \( b = 5 \) for bootstrapping the residuals.

\(^{18}\)Fund #154, Paradigm Master Fund, reported one unusually large negative return, see Figure 2. As a result, it unduly dominates the bootstrap sampling distribution of the largest studentized test statistics \( z_{120,r_1} \), see Figure 3.
only. The \(p\)-values are obtained by the studentized circular block bootstrap, which corresponds to applying the StepM method to each single strategy, that is, the special case \(S = 1\). The block sizes for the circular block bootstrap are chosen, individually for each fund, via Algorithm A.5 for the special case \(S = 1\). The semi-parametric model \(\tilde{P}_T\) used in this algorithm is an AR(1) model in conjunction with bootstrapping the residuals.\(^{19}\) The results are displayed in the left half of Table 3.

Next, we turn to the studentized StepM method and its generalizations.\(^{20}\) The block sizes for the circular block bootstrap are chosen via Algorithm A.5. The semi-parametric model \(\tilde{P}_T\) used in this algorithm is a VAR(1) model in conjunction with bootstrapping the residuals.\(^{21}\) The \(k\)-StepM procedures are based on the operative method using \(N_{\text{max}} = 100\); see Remark 4.2. The results are displayed in the right half of Table 3.

Not surprisingly, the results are comparable to those of the simulation study. First, by controlling a generalized error rate, the number of rejected hypotheses can greatly increase. For example, for the nominal level of \(\alpha = 0.1\), while the (1-)StepM method rejects 16 hypotheses, the 2-StepM method rejects 29 hypotheses. Second, the methods that implicitly account for the dependence structure of the test statistics reject more hypotheses than the methods based on individual \(p\)-values. For example, for the nominal level of \(\alpha = 0.1\), while the FDP-LR\(_{0.1}\) method rejects 22 hypotheses, the FDP-StepM\(_{0.1}\) method rejects 36 hypotheses. Third, the methods controlling a central tendency of the FDP are the ones which reject the most hypotheses.

### 9.2 Multiple Regression

In empirical work, it is quite common to estimate large-dimensional regression models and to then ask which are the ‘important’ variables. The habitual practice is to assess ‘importance’ via the individual \(t\)-statistics or, equivalently, via the individual \(p\)-values without taking into account the multitude of tests. Consequently, as discussed earlier, typically too many variables will be identified as ‘important’.

As an example, we consider a Mincer regression where log wages are regressed on a large number of explanatory variables. The data consists of a random sample of \(n = 4,975\) persons from the Austrian Social Security data base on 08/10/2001. The explanatory variables include a dummy for gender, a dummy for blue collar (vs. white collar), age, age squared, work experience, work experience squared, time at current company, time at current company squared, state dummies,

\(^{19}\)To account for leftover dependence not captured by the AR(1) model, we use the stationary bootstrap with average block size \(b = 5\) for bootstrapping the residuals.

\(^{20}\)Similar to the analysis of Romano and Wolf (2005c), the basic StepM method does not detect a single outperforming fund, so it is not pursued further.

\(^{21}\)To account for leftover dependence not captured by the VAR(1) model, we use the stationary bootstrap with average block size \(b = 5\) for bootstrapping the residuals.
industry dummies, and state-industry interaction dummies, as well as an intercept. The total number of explanatory variables is $S = 291$.

We now use the various multiple testing methods to identify the 'important' variables, starting with the Holm procedure and its generalizations as well as the FDR procedure of Benjamini and Hochberg (1995), all of which are based on individual $p$-values only. The $p$-values are obtained by the wild bootstrap to account for possible heteroskedasticity. To generate the resampled errors, we use the two-point distribution; see (6.21) in Hinkley and Davison (1997). Standard errors both in the real world and in the bootstrap world are computed via the well-known White estimator. The White estimator uses the modified residuals rather than the raw residuals, since the former have equal variance; see page 271 in Hinkley and Davison (1997). The results are displayed in the left half of Table 4.

Next, we turn to the studentized StepM method and its generalizations. The $k$-StepM procedures are based on the operative method using $N_{\text{max}} = 100$; see Remark 4.2. The results are displayed in the right half of Table 4.

10 Conclusions

The problem of testing multiple hypotheses is ubiquitous in econometric applications. Unfortunately, this problem is very often simply ignored. As a result, too many true null hypotheses will be rejected. The classical approach to account for the multitude of hypotheses under test is to control the familywise error rate (FWE), defined as the probability of falsely rejecting even one true hypothesis. But when the number of hypotheses is very large, this criterion can become too stringent. As a result, potentially very few false hypotheses will be rejected.

This paper has reviewed various generalized error rates. They are more liberal than the FWE yet still account for the multitude of tests by allowing for a small number or a small (expected) proportion of true hypotheses among all rejected hypotheses. Some simulations as well as two empirical applications have demonstrated that in this way many more false hypotheses can be rejected compared to control of the FWE.

As a special emphasis, we have presented some very recent multiple testing procedures that implicitly account for the dependence structure of the individual test statistics via an application of the bootstrap. The advantage over traditional multiple testing procedures based on individual $p$-values alone is that the number of false hypotheses rejected often increases, while the control of the generalized error rates is not sacrificed. This advantage has also been highlighted via simulations
and two empirical applications. The disadvantage is the increased computational cost, but due to the availability of fast computers this is less and less of a concern.

We have discussed further how the control of generalized error rates can apply to various notions of model selection.
A Use of the Bootstrap

This appendix details how to compute the constants $\hat{c}_j$, $\hat{c}_{j,|\cdot|}$, $\hat{d}_j$, and $\hat{d}_{j,|\cdot|}$ in Algorithms 4.1, 4.2, 4.3, and 4.4, respectively, via the bootstrap. At first, a proper choice of the estimator $\hat{P}_T$ of the underlying probability mechanism $P$ must be made. (One can implicitly define $\hat{P}_T$ by describing a how a bootstrap data matrix $X_T^*$ is generated from $\hat{P}_T$.) This choice depends on the context. If the data $X_{1}^{(T)}, \ldots, X_{T}^{(T)}$ are i.i.d., one should choose Efron’s (1979) bootstrap; if they constitute a time series, one should choose the moving blocks bootstrap, the circular blocks bootstrap, or the stationary bootstrap. These various bootstrap methods are detailed in Appendix B of Romano and Wolf (2005c). In any case, we use the notation $\hat{\theta}_T$ for a suitable parameter vector corresponding to the bootstrap law.

Algorithm A.1 (Computation of the $\hat{c}_j$ via the Bootstrap)

1. The labels $r_1, \ldots, r_S$ and the numerical values of $R_1, R_2, \ldots$ are given in Algorithm 4.1.

2. Generate $M$ bootstrap data matrices $X_{T,1}^*, \ldots, X_{T,M}^*$. (One should use $M \geq 1,000$ in practice.)

3. From each bootstrap data matrix $X_{T,m}^*$, $1 \leq m \leq M$, compute the individual test statistics $w_{T,1,m}^*, \ldots, w_{T,S,m}^*$.

4. (a) For $1 \leq m \leq M$, and any needed $K$, compute $kmax_{T,K}^* = k\cdot\max_{s \in K}(w_{T,r_s}^* - \hat{\theta}_{T,r_s})$.

   (b) Compute $c_K(1 - \alpha, k, \hat{P}_T)$ as the $1 - \alpha$ empirical quantile of the $M$ values $kmax_{T,K}^{*,1}, \ldots, kmax_{T,K}^{*,M}$.

5. If $j = 1$, $\hat{c}_1 = c_{\{1, \ldots, S\}}(1 - \alpha, k, \hat{P}_T)$

   If $j > 1$, $\hat{c}_j = \max\{c_K(1 - \alpha, k, \hat{P}_T) : K = I \cup \{R_{j-1} + 1, \ldots, S\}, I \subset \{1, \ldots, R_{j-1}\}, |I| = k - 1\}$

Algorithm A.2 (Computation of the $\hat{c}_{j,|\cdot|}$ via the Bootstrap)

1. The labels $r_1, \ldots, r_S$ and the numerical values of $R_1, R_2, \ldots$ are given in Algorithm 4.2.

2. Generate $M$ bootstrap data matrices $X_{T,1}^*, \ldots, X_{T,M}^*$. (One should use $M \geq 1,000$ in practice.)

3. From each bootstrap data matrix $X_{T,m}^*$, $1 \leq m \leq M$, compute the individual test statistics $w_{T,1,m}^*, \ldots, w_{T,S,m}^*$.

4. (a) For $1 \leq m \leq M$, and any needed $K$, compute $kmax_{T,K,|\cdot|}^* = k\cdot\max_{s \in K}|w_{T,r_s}^* - \hat{\theta}_{T,r_s}|$. 

32
(b) Compute $c_{k_{j_{||}}}(1 - \alpha, k, \hat{P}_T)$ as the $1 - \alpha$ empirical quantile of the $M$ values $\text{kmax}_{T,K_{||}}^{*,1}, \ldots, \text{kmax}_{T,K_{||}}^{*,M}$.

5. If $j = 1$, $\hat{d}_{1_{||}} = c_{1_{||}}(1 - \alpha, k, \hat{P}_T)$
If $j > 1$, $\hat{d}_{j_{||}} = \max\{c_{k_{j_{||}}}(1 - \alpha, k, \hat{P}_T) : K = I \cup \{R_{j-1} + 1, \ldots, S\}, I \subset \{1, \ldots, R_{j-1}\}, |I| = k - 1\}

Algorithm A.3 (Computation of the $\hat{d}_j$ via the Bootstrap)

1. The labels $r_1, \ldots, r_S$ and the numerical values of $R_1, R_2, \ldots$ are given in Algorithm 4.3.

2. Generate $M$ bootstrap data matrices $X^*_T^{*,1}, \ldots, X^*_T^{*,M}$. (One should use $M \geq 1,000$ in practice.)

3. From each bootstrap data matrix $X^*_T^{*,m}$, $1 \leq m \leq M$, compute the individual test statistics $w^*_{T,1}^{*,m}, \ldots, w^*_{T,S}^{*,m}$. Also, compute the corresponding standard errors $\hat{\sigma}^*_{T,1}^{*,m}, \ldots, \hat{\sigma}^*_{T,S}^{*,m}$.

4. (a) For $1 \leq m \leq M$, and any needed $K$, compute $\text{kmax}_{T,K_{||}}^{*,m} = \text{k-max}_{s \in K}(\{|w^*_{T,r_s} - \hat{\theta}_{T,r_s}|/\hat{\sigma}^*_{T,r_s}^{*,m}\}$.
(b) Compute $d_K(1 - \alpha, k, \hat{P}_T)$ as the $1 - \alpha$ empirical quantile of the $M$ values $\text{kmax}_{T,K_{||}}^{*,1}, \ldots, \text{kmax}_{T,K_{||}}^{*,M}$.

5. If $j = 1$, $\hat{d}_{1_{||}} = d_{1_{||}}(1 - \alpha, k, \hat{P}_T)$
If $j > 1$, $\hat{d}_{j_{||}} = \max\{d_K(1 - \alpha, k, \hat{P}_T) : K = I \cup \{R_{j-1} + 1, \ldots, S\}, I \subset \{1, \ldots, R_{j-1}\}, |I| = k - 1\}$

Algorithm A.4 (Computation of the $\hat{d}_{j_{||}}$ via the Bootstrap)

1. The labels $r_1, \ldots, r_S$ and the numerical values of $R_1, R_2, \ldots$ are given in Algorithm 4.4.

2. Generate $M$ bootstrap data matrices $X^*_T^{*,1}, \ldots, X^*_T^{*,M}$. (One should use $M \geq 1,000$ in practice.)

3. From each bootstrap data matrix $X^*_T^{*,m}$, $1 \leq m \leq M$, compute the individual test statistics $w^*_{T,1}^{*,m}, \ldots, w^*_{T,S}^{*,m}$. Also, compute the corresponding standard errors $\hat{\sigma}^*_{T,1}^{*,m}, \ldots, \hat{\sigma}^*_{T,S}^{*,m}$.

4. (a) For $1 \leq m \leq M$, and any needed $K$, compute $\text{kmax}_{T,K_{||}}^{*,m} = \text{k-max}_{s \in K}(\{|w^*_{T,r_s} - \hat{\theta}_{T,r_s}|/\hat{\sigma}^*_{T,r_s}^{*,m}\}$.
(b) Compute $d_K(1 - \alpha, k, \hat{P}_T)$ as the $1 - \alpha$ empirical quantile of the $M$ values $\text{kmax}_{T,K_{||}}^{*,1}, \ldots, \text{kmax}_{T,K_{||}}^{*,M}$.

5. If $j = 1$, $\hat{d}_{1_{||}} = d_{1_{||}}(1 - \alpha, k, \hat{P}_T)$
If $j > 1$, $\hat{d}_{j_{||}} = \max\{d_K(1 - \alpha, k, \hat{P}_T) : K = I \cup \{R_{j-1} + 1, \ldots, S\}, I \subset \{1, \ldots, R_{j-1}\}, |I| = k - 1\}$
Remark A.1 For convenience, one can typically use $w_{T,r_k}$ in place of $\hat{\theta}_{T,r_k}$ in step 4(a) of Algorithms A.1–A.4. Indeed, the two quantities are the same under the following conditions: (1) $w_{T,k}$ is a linear statistic; (2) $\theta_k = \text{E}(w_{T,k})$; and (3) $\hat{\theta}_T$ is based on Efron’s bootstrap, the circular blocks bootstrap, or the stationary bootstrap. Even if conditions (1) and (2) are met, $w_{T,r_k}$ and $\theta_{T,r_k}$ are not the same if $\hat{\theta}_T$ is based on the moving blocks bootstrap due to ‘edge’ effects; see Appendix B of Romano and Wolf (2005c). On the other hand, the substitution of $w_{T,r_k}$ for $\hat{\theta}_{T,r_k}$ does not affect in general the consistency of the bootstrap approximation. Lahiri (1992) discusses this subtle point for the special case of time series data and $w_{T,r_k}$ being the sample mean. He shows that centering by $\hat{\theta}_{T,r_k}$ provides second-order refinements but is not necessary for first-order consistency.

When a time series bootstrap is used, then the choice of the (average) block size becomes an important practical problem. The method we propose here to choose a block size for an application of the $k$-StepM procedure is a generalization of Algorithm 7.1 of Romano and Wolf (2005c) who only deal with the StepM procedure.

Consider the first step of the $k$-StepM procedure. The goal is to construct a generalized joint confidence region for the parameter vector $\theta$ with nominal coverage probability of $1 - \alpha$. Here, importantly, ‘coverage probability’ stands for the probability of containing at least $S - k + 1$ elements of $\theta$.

Algorithm A.5 (Choice of Block Size)

1. Fit a semiparametric model $\hat{P}_T$ to the observed data $X_T$.
2. Fix a selection of reasonable block sizes $b$.
3. Generate $M$ data sets $\tilde{X}_T^{1}, \ldots, \tilde{X}_T^{M}$ according to $\hat{P}_T$.
4. For each data set $\tilde{X}_T^{m}$, $m = 1, \ldots, M$, and for each block size $b$, compute a generalized joint confidence region $GJCR_{m,b}$ for $\theta$.
5. Compute $\hat{g}(b) = \#\{\text{At least } S - k + 1 \text{ elements of } \theta(\hat{P}_T) \in GJCR_{m,b}\}/M$.
6. Find the value of $\bar{b}$ that minimizes $|\hat{g}(b) - (1 - \alpha)|$ and use this value $\bar{b}$.

Algorithm A.5 is based on the first step of the $k$-StepM method. Since the general $k$-StepM method, for $k > 1$, does not discard any hypotheses in subsequent steps — in contrast to the StepM method — we recommend to keep using the chosen value $\bar{b}$ throughout. If, on the other hand, the operative method of Remark 4.2 is used, then at a given subsequent step some hypotheses may already have been discarded. In that case, one can apply Algorithm A.5 to the subset of $\theta$ which corresponds to the non-discarded hypotheses.
References


37


Table 1: Empirical FWEs, FDPs, and FDRs (in the rows ‘Control’ and expressed as percent) and average number of false hypotheses rejected (in the rows 'Rejected') for various methods, with $T = 100$ and $S = 500$. The nominal level is $\alpha = 5\%$, apart from the second to last column where it is $\alpha = 50\%$. The number of repetitions is 5,000 when all $\theta_s = 0$ and 2,000 for all other scenarios; and the number of bootstrap resamples is $M = 200$.

<table>
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<tr>
<th></th>
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<th>Common correlation: $\rho = 0.5$</th>
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<td>10-gH</td>
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<tr>
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Table 2: The ten largest basic and studentized test statistics, together with the corresponding hedge funds, in our empirical application. The return unit is 1 percent.

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<th>Fund</th>
<th>(z_{T,s})</th>
<th>Fund</th>
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<td>1.34</td>
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<tr>
<td>1.31</td>
<td>Spinner Global Technology</td>
<td>6.32</td>
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Table 3: Number of outperforming funds identified.

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Table 4: Number of important variables identified.

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Figure 1: Boxplots of realized FDPs for various scenarios. The upper part is for control of the median FDP while the lower part is for control of the FDR. The labels on the x-axis — "100", "200", and "400" — denote the number of false hypotheses. The horizontal dashed line indicates $\gamma = 0.1$. 
Figure 2: Histogram of the monthly log returns of fund #154. In 08/1995 the fund, Paradigm Master Fund, reported a return of $-53.77\%$, resulting in a tremendous outlier to the left.
Figure 3: Histogram of the fund index that corresponds to the largest studentized statistic $z_{120, r_1}^*$ in $M = 5,000$ bootstrap repetitions. Fund #154, Paradigm Master Fund, corresponds to the largest studentized statistic disproportionately often.