NONPARAMETRIC BAYESIAN MULTIPLE HYPOTHESIS TESTING OF AUTOREGRESSIVE TIME SERIES

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This paper describes a framework for flexible multiple hypothesis testing of autoregressive time series. The modeling approach is Bayesian, thought a blend of frequentist and Bayesian reasoning is used to evaluate procedures. Nonparametric characterizations of both the null and alternative hypotheses will be shown to be the key robustification step necessary to ensure reasonable Type-I error performance. The methodology is applied to part of a large database containing up to 50 years of corporate performance statistics on 24,157 publicly traded American companies, where the primary goal of the analysis is to flag companies whose historical performance is significantly different from that expected due to chance.

1. Introduction.

1.1. Multiple testing of time series. Suppose a single time series of length $T$ is observed for each of $N$ different units. Two possible models for each time series are entertained: a simple autoregressive null model $M_0$ and a more complex alternative model $M_A$. The goal is to determine which units come from the alternative model.

This is a frequent problem in the analysis of multiple time series, and although many details will be context-dependent, certain common themes emerge. Of key interest is how, in repeatedly applying a procedure used for testing a single time series, the rate of Type-I errors can be controlled. Model-based approaches are an attractive option, but model errors can become overwhelming in the face of massive multiplicity. One of this paper’s main results is that great care must be taken in characterizing $M_0$ and $M_A$ in order to keep false positives at bay, with the suggested robustification step involving the use of nonparametric Bayesian methods.

*This research was supported by a graduate research fellowship from the U.S. National Science Foundation. The author would also like to thank: Mumtaz Ahmed, Michael Raynor, Lige Shao, Jim Guszcza, and Jim Wappler of Deloitte Consulting for their insight into the application discussed here; along with Andy Henderson of the University of Texas and Carlos Carvalho of the University of Chicago for all their helpful feedback.

AMS 2000 subject classifications: Primary 62G10, 62M02; secondary 62P20, 62G35

Keywords and phrases: Multiple testing, Bayesian model selection, nonparametric Bayes, financial time series
In typical hypothesis-testing scenarios involving standard parametric models $M_0$ and $M_A$, the Bayes factor $BF(M_A : M_0)$ contains an “Ockham’s razor” term (Jefferys and Berger, 1992) that penalizes the more complex model. In most cases this is the result of needing to integrate the likelihood across a higher-dimensional prior under the more complex model, which will therefore have a more diffuse predictive distribution. This penalty for model complexity is quite different from any sort of penalty term imposed for conducting multiple hypothesis tests (Scott and Berger, 2008).

But the multiple-testing framework employed here uses Dirichlet-process mixtures to represent the unknown null and alternative models, which are properly thought of as being infinite-dimensional. Another of this paper’s main objectives is to clarify the nature and operating characteristics of this penalty for model complexity in nonparametric settings—specifically, how it interfaces with the recommended multiple-testing methodology.

1.2. Motivating example. A running example from management theory will be used to motivate and study the proposed Bayesian model on a real data set comprising up to 50 years of annual performance information for over 24,000 publicly traded American companies. The goal of the analysis is to flag firms whose performance is highly unlikely to have occurred by random chance, since these firms may have good (or bad) management practices that are discernible through follow-up case studies. The results and discussion are problem-specific, but areas in which the methodology and lessons can be generalized will be pointed out.

Longitudinal performance stratification is a classic topic in management theory. Indeed, one of the primary aims of strategic-management research (and the conceit of many best-selling books) is to explain why some firms fail and others succeed.

Much academic work in this direction focuses on decomposing observed variation into market-level, industry-level, and firm-level components (Bowman and Helfat, 2001; Hawawini et al., 2003). Of the work that attempts to identify specific non-random performers, much of it relies upon model-free clustering algorithms (for example, Harrigan, 1985), which contain no guarantee that the clusters found will be significantly different from one another. Other approaches employ simple classical tests (Ruefli and Wiggins, 2000), often based upon ordinal time series. These have the advantage of being model-free, but are typically not based upon sufficient statistics, and suffer from the fact that available multiplicity-correction approaches (e.g. Bonferroni correction) tend toward the overly aggressive.

Due to the number of firms for which public financial data is available,
this problem makes an excellent testbed for the study of general-purpose multiple-testing methodology in time-series analysis. There is, however, no theoretical ideal of what an “average-performing” company should look like, beyond the notion that it should revert to the population-level mean even if it has some randomly good or bad years. The Bayesian approach requires that suitable notions of randomness and nonrandomness be embodied in a statistical model. This model must confront an obvious multiplicity problem: many thousands of companies will be tested, and false positives will make expensive wild-goose chases out of any follow-up studies seeking to explain possible sources of competitive advantage. Robustness and trustworthy Type-I error characteristics are therefore crucial practical considerations, and so even though this paper’s modeling approach is Bayesian, it also contains much frequentist reasoning regarding Type-I error rates.

2. Testing a zero-mean AR(1) model.

2.1. The model. This section outlines a basic framework for multiple testing that, for the sake of illustration, will be purposely simplistic. Nonetheless, it will provide a useful jumping-off point for the methodological developments of subsequent sections, and will show why more flexible models are typically needed in order to achieve reasonable Type-I error performance.

Let $y_{it}$ be the observation for unit $i$ at time $t$, and let $y_i$ be the vector of observations for unit $i$. In the management-theory example, $y$ is standardized performance metric called Return on Assets (ROA), which measures how efficiently a company’s assets generate earnings. Each company’s ROA values were regressed upon a set of covariates judged to be relevant by three experts in management theory collaborating on the project. These include the company’s size, debt-to-equity ratio, and market share, along with categorical variables for year and for industry membership. (See Ruefli and Wiggins, 2002, for a summary of the literature regarding covariate effects on observed firm performance.) The actual values used in the following analyses were the residuals from this regression. Also, since the question at issue is one of relative performance, not absolute performance, these residuals were standardized by CDF transform to follow a $N(0,1)$ distribution.

Since we do not expect random gains or losses in one year to be completely erased by the following year, a model accounting for serial autocorrelation seems mandatory. Management-theoretic support for this assumption in the present context can be found in Denrell (2003) and Denrell (2005); analogous situations in engineering, finance, and biology are very common.

The null hypothesis is then a stationary AR(1) model depending upon
parameter $\param = (\phi, \nu)$:

$$y_{it} = \phi y_{i,(t-1)} + \nu_t$$
$$\nu_t \overset{iid}{\sim} N(0, \nu) .$$

This assumption allows for long runs of good or bad performance due simply to chance: a large shock ($\nu_t$) may take quite awhile to decay depending upon the value of $\phi$, which is assumed to lie on $(-1, 1)$.

Non-null companies can then be modeled as AR(1) processes that revert to a nonzero mean. Placing a mixture prior on this unknown mean will then encode the relevant hypothesis test:

$$y_i \sim N(y_i \mid m_i 1, \Sigma_{\theta}) \quad (1)$$
$$\theta \sim N(\phi \mid d, D) \times IG(\nu \mid a, b) \quad (2)$$
$$m_i \sim p \cdot N(0, \sigma^2) + (1 - p) \delta_0 \quad (3)$$

with $1$ is the vector of all ones, $\Sigma_{\theta}$ is the familiar AR(1) variance matrix, $\delta_0$ is a point mass at 0, and $p \in [0, 1]$ is the prior probability of arising from the alternative hypothesis. The exchangeable normal prior on the nonzero means $m_i$ reflects the prior belief that, among firms that systematically deviate from zero, most of the deviations will be relatively small.

The posterior probabilities $p_i = P(m_i \neq 0 \mid Y)$ then can be used to flag non-null units. In some contexts, these are called the posterior inclusion probabilities, reflecting inclusion in the “non-null” set. One could simply choose a small value of $p$, with the expectation that a prior bias toward the null will solve the multiplicity problem. But the key intuition in using this model for multiple hypothesis testing is to treat $p$ as just another model parameter to be estimated from the data, giving it a uniform prior. This induces an effect that is often referred to as an automatic multiple-testing penalty, with the effect being “automatic” in the sense that no arbitrary penalty terms must be specified.

This effect can most easily be seen if one imagines repeatedly testing a fixed number of signals in the presence of an increasing number of null units. As “noise” comes to dominate the cohort of tested units, the posterior mass of $p$ will concentrate near 0, making it increasingly difficult for all units (even the signals) to overcome the prior belief in their irrelevance. This yields much the same effect as choosing a small value for $p$ after the fact, but Bayesian learning about $p$ negates the need to make such an arbitrary choice.

It is important to distinguish this fully Bayesian model from the empirical-Bayes approach to multiplicity correction, whereby $p$ is estimated by maximum likelihood. These two approaches are intuitively similar, since the prior
inclusion probability is estimated by the data in order to yield an automatic multiple-testing penalty. Yet the approaches have quite different operational and theoretical properties (Scott and Berger, 2008), with the focus here being on the fully Bayesian approach.

Similar multiple-testing procedures have been extensively studied in many different contexts. See Scott and Berger (2006) for theoretical development and for an example involving DNA microarrays; Carvalho and Scott (2007) for portfolio selection; Carvalho et al. (2008) for cancer genomics; and George and Foster (2000) and Cui and George (2008) for examples in regression.

The model must be completed by specifying priors for $\phi$, $v$, and $\sigma^2$. In the example at hand, the data have been standardized to a $N(0, 1)$ scale, so it makes sense to choose $\sigma^2 = 1$. In more general settings, the prior for $\sigma^2$ must be appropriately scaled by $\phi$ and $v$ in the absence of strong prior information, since the marginal variance of the residual autoregressive process is the only quantity that provides a default scale for the problem.

The conditional likelihoods of each data vector under the two hypotheses ($m_i \neq 0$ versus $m_i = 0$) are available in closed form:

\begin{align}
P(y_i | m_i = 0, \phi, v) & = N\left(y_i | 0, \frac{v}{1 - \phi^2} \Phi\right) \\
P(y_i | m_i \neq 0, \phi, v) & = N\left(y_i | 0, \frac{v}{1 - \phi^2} \Phi + \sigma^2(11^t)\right)
\end{align}

where $\Phi$ is the standard AR(1) correlation matrix and where $(11^t)$ is the matrix of all ones.

The ratio of (5) to (4) gives the Bayes factor, conditional upon $\phi$ and $v$, for testing an individual time series against the null model. Taken as a group, these Bayes factors provide information about the unknown prior inclusion probability $p$; a preponderance of large Bayes factors (suggesting many non-null units in the study) will cause $p$ to be large, and vice versa.

The posterior inclusion probabilities $\{p_i\}$ can be computed straightforwardly using importance sampling to account for posterior uncertainty about $\phi$, $v$, and $p$, which will be stable as long as $p$ is not too close to 0 or 1. The advantage of importance sampling here is that a common importance function may be used for all marginal inclusion probabilities, greatly simplifying the required calculations. After transforming all variables to have unrestricted domains, importance sampling was used to compute the results presented in this section, with repetition and plots of the importance weights used to confirm stability.

### 2.2. Results on ROA data

Historical ROA time series for 3,459 publicly traded American companies between 1965 and 2004 were used to fit the
above model. This encompasses almost every public company over that pe-
period for which at least 20 years of data were available. Standard independent
conjugate priors for $\phi$ and $v$ were used:

\begin{align}
\phi &\sim N_U(0.5, 0.25^2) \\
v &\sim IG(2, 1),
\end{align}

where $N_U$ indicates that the normal prior for $\phi$ is truncated to lie on the
appropriate interval. These priors were chosen to reflect the expectations of
the collaborating management theorists regarding the persistence and scale
of random ROA fluctuations from year to year. Because these parameters
appear in both the null and alternative models, integrals over these priors
appear in both the numerator and denominator of the Bayes factor com-
paring $m_i \neq 0$ versus $m_i = 0$, making them not overly influential in the
analysis; see Berger et al. (1998) for a general guidelines on choosing com-
mon hyperparameters in model-selection problems.

In many cases the results of the fit seemed reasonable. Most firms were
assigned to $M_0$ with high probability, suggesting strong multiplicity control,
while companies with obvious patterns of sustained excellence or inferior-
ity were flagged as being from $M_A$ with very high probability. Figure 1
contains instructive examples: two excellent companies (WD-40 and Coca-
Cola), along with one obviously poor company (Oglethorpe Power) were
assigned greater than 95% probability of being non-null. A fourth example,
Texas Intruments, had several intermittent years of good performance but
no pattern of sustained excellence, and the model gave it greater than 90%
probability of being from the null model.

On the other hand, the model displayed two serious shortcomings:

- Many firms diverged in obvious ways (for example, via the appearance
  of long-term trends) from the expectations of a single AR(1) model.
  Two such examples are in Figure 2. Discussion of this important issue
  is postponed until Section 4.
- More subtly, the model imposed a homogeneous error structure on
data that seemed rather heterogeneous. Some fairly basic exploratory
data analysis indicated that firms displayed differing degrees of “stick-
iness” in their trajectories. This suggested that a single value of $\phi$
for the entire data set might be unsatisfactory. Likewise, some firms
appeared systematically more volatile than others, making a single-
variance model equally questionable.

2.3. Robustness simulations. The possibility of model errors in (1)–(3)
bring the issue of robustness to the forefront. This section describes the
**Fig 1.** Four example companies.

**Fig 2.** Two examples that seem to display trends.
results of a simulation study that shows just how poorly this model can perform when a particular type of model error is encountered: deviation from the “single $\phi$, single $v$” approach to describing the AR(1) residuals of all companies in the sample.

Several data sets displaying different levels of heterogeneity were simulated. The homogeneous (i.e. single $\phi$, single $v$) model was subsequently fit to each simulated data set in order to assess the robustness of the procedure’s Type-I error performance.

Each simulated data set had 3500 times series of length $T = 40$, with each time series drawn from a mixture distribution of AR(1) models. These distributions ranged from trivial one-component mixtures (for which the assumed model was true) to complex nine-component mixtures (for which the assumed model was quite a bad approximation). These conditions are summarized in Table 1. In the four- and nine-component models, all components were equiprobable. Since all simulated companies had $m_i = 0$, ideally there should be no positive flags.

For the purposes of classification, thresholding is reported at the $p_i \geq 0.5$ and the $p_i \geq 0.9$ levels, where $p_i$ is the posterior inclusion probability for company $i$. The first ($p_i \geq 0.5$) threshold reflects a 0–1 loss function that symmetrically penalizes false positives and false negatives. The second threshold is arbitrary, but meant to reflect a more conservative approach to identifying signals. A full decision-theoretic analysis incorporating more realistic loss functions would yield a different, data-adaptive threshold.

Table 1 supports two conclusions:

- The proposed model yields very strong control over false positives when its assumptions are met: 3 false positives and 0 false positives in the two cases investigated, out of 3,500 units tested. This confirms that the theory outlined in Scott and Berger (2006), which concerns the much simpler normal-means testing problem, applies here as well.
- This excellent Type-I error profile is not at all robust to a violation of the autoregressive model’s assumptions. In the most extreme case, nearly a third of units (1045 out of 3500) tested had inclusion probabilities $p_i \geq 50\%$, when in reality none were from the alternative model. In other less extreme cases, the false positives still numbered in the hundreds, which is clearly unsatisfactory.

These results dramatically illustrate the effect of heterogeneity in the autoregressive profiles of each tested unit. If such heterogeneity exists but is ignored, the Type-I error performance of the procedure may be severely compromised.
3. A nonparametric null model. In the previous section, a specific form of model error—different groups of companies following different AR models—was shown to be a source of overwhelming Type-I errors. Hence a natural extension is to consider a more complicated autoregressive model for the residuals that accounts for the possibility of stratification.

The Dirichlet process (Ferguson, 1973) offers a straightforward nonparametric technique for accommodating uncertainty about this random distribution. Let \( z_i \) represent the response vector for unit \( i \), for now ignoring any contribution due to a nonzero mean. Recall that for parameter \( \theta = (\phi, v) \), \( \Sigma_{\theta} \) denotes the AR(1) variance matrix. The DP mixture model can then be written as a hierarchical model:

\[
\begin{align*}
\mathbf{z}_i & \sim \mathcal{N}(\mathbf{z}_i \mid \mathbf{0}, \Sigma_{\theta_i}) \\
\theta_i & \sim \mathcal{G}, \quad \mathcal{G} \sim \text{DP}(\alpha, G_0) \\
G_0 & = \mathcal{N}(\phi \mid d, D) \times \text{IG}(v \mid a, b),
\end{align*}
\]

where the hyperparameters \((d, D)\) and \((a, b)\) must be chosen to reflect the expected properties of the base measure \( G_0 \) (which is a product of two independent distributions, a normal and an inverse-gamma), and where \( \alpha \) controls the degree of expected departure from the base measure.

Dirichlet-process priors for nonparametric Bayesian density estimation were popularized by Escobar and West (1995), and their use in nonlinear autoregressive time series dates to Müller et al. (1997). For another reference on the use of DP priors in the context of Bayesian multiple comparisons, see Gopalan and Berry (1998).

Realizations of the Dirichlet process are almost surely discrete distributions, and so we expect some of the \( \theta_i \)'s to be the same across companies. This is the DP framework’s primary strength here, since it will facilitate borrowing of information across time series. Simply allowing each time series

<table>
<thead>
<tr>
<th>Number of Components: Model</th>
<th>( \hat{p} )</th>
<th>( # p_i \geq 0.5 )</th>
<th>( # p_i \geq 0.9 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1: ((\phi, v) = (0.5, 0.25))</td>
<td>0.01</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>1: ((\phi, v) = (0.9, 0.5))</td>
<td>0.02</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>4: ((\phi, v) \in {0.5, 0.7} \times {0.25, 0.5})</td>
<td>0.02</td>
<td>30</td>
<td>4</td>
</tr>
<tr>
<td>4: ((\phi, v) \in {0.4, 0.6, 0.8} \times {0.05, 0.25, 0.5})</td>
<td>0.08</td>
<td>152</td>
<td>66</td>
</tr>
<tr>
<td>9: ((\phi, v) \in {0.2, 0.95} \times {0.05, 0.5})</td>
<td>0.37</td>
<td>1045</td>
<td>560</td>
</tr>
<tr>
<td>9: ((\phi, v) \in {0.2, 0.6, 0.95} \times {0.05, 0.25, 0.5})</td>
<td>0.29</td>
<td>797</td>
<td>493</td>
</tr>
</tbody>
</table>

Table 1

Robustness of the multiple-testing procedure’s Type-I error performance to heterogeneity in the autoregressive profiles of tested units. Here \( \hat{p} \) refers to the posterior mode of the mixing ratio \( p \), and the \( p_i \)'s are the posterior probabilities that each \( m_i \neq 0 \).
to have its own $\phi$ and own $v$ would make for a simpler model (albeit with more parameters), but the DP prior reflects the subject-specific knowledge that significant clustering of autoregressive parameters should be expected.

This will lead to behavior similar to that predicted by a finite mixture of AR(1) models, such as the kind considered by Frühwirth-Schnatter and Kaufmann (2008). The Dirichlet-process prior, however, avoids the complicated task of directly computing marginal likelihoods for mixture models of different sizes, and so makes computation much simpler. Note that since the marginal distribution of one draw from a Dirichlet-process mixture depends only on the base measure, the DP acts like a mixture model that is predictively matched to a single observation.

It is important to consider, of course, how choices for $\alpha$ and $G_0$ affect the implied prior distributions both for the number of mixture components and for the parameters associated with each component. The marginal prior for the parameters of each mixture component is simply given by the base measure, while for certain choices of $\alpha$ the number of mixture components $k$ is asymptotically Poisson: if $\alpha = \lambda / \log n$, then $k$ is approximately distributed as $1 + \text{Po}(\lambda)$ as $n \to \infty$ (Dunson and Herring, 2006).

4. A nonparametric alternative model.

4.1. Trajectories as random functions. Section 2 considered a simple constant-mean AR(1) model for non-null units, and Section 3 modified the AR(1) assumption to account for a richer autoregressive structure. This section now modifies the constant-mean assumption to allow for time-varying nonzero trajectories upon which the autoregressive residuals are superimposed. Most management teams, after all, do not stay the same for 40 or 50 years, and we should not expect their performance to stay the same, either.

Firm performance trajectories can be viewed as continuous random functions that are observed at discrete (in this case, annual) intervals. This is essentially a nonparametric version of a mixed-effects model for longitudinal data. Similar models have been considered by Bigelow and Dunson (2005), Dunson and Herring (2006), and, in a spatial context, Gelfand et al. (2005).

Let $\mathbf{f}_i = \{y_i(t), t \in \mathbb{R}^+\}$ be a continuous-time stochastic process for each observed unit, and let $\mathbf{t}_i$ denote the vector of times at which each unit was observed. Then the model is

\begin{align}
\mathbf{y}_i &= \mathbf{f}_i(\mathbf{t}_i) + \mathbf{z}_i \\
\mathbf{z}_i &\sim \mathcal{N}(\mathbf{z}_i | \mathbf{0}, \Sigma_{\theta_i}), \quad \mathbf{\theta}_i \sim G \\
\mathbf{f}_i &\sim p \cdot \mathcal{F} + (1 - p) \cdot \delta_{F_0}
\end{align}
where $G$ is the nonparametric residual model defined in Section 3, $\delta_{F_0}$ represents a point mass at the zero function $F_0(t) = 0$, and $F$ is a random distribution over a function space $\Omega$. As in Section 2, $p$ is the unknown prior probability of coming from the alternative model $M_A$, represented in this case by the distribution $F$. It is convenient to represent each hypothesis test using a model index parameter $\gamma_i$: $\gamma_i = 0$ if $f_i = F_0$ (i.e. the null model $M_0$ is true for unit $i$), and $\gamma_i = 1$ otherwise.

4.2. Choice of features for $M_A$. The crucial consideration in using the above model for hypothesis testing is that the space $\Omega$ from which each $f_i$ is drawn must be restricted to a sufficiently small class of functions. This would be necessary even if $F$ were only being estimated, and not tested against a simpler model: if $\Omega$ is too broad, then the alternative model itself will not be likelihood-identified, since any pattern of residuals could equally well be absorbed by the mean function.

But this guideline is even more important in model-selection problems; an over-broad class of functions will mean that the random distribution $F$ is vague, in the sense that the predictive distribution of observables will be diffuse. It is widely known that using vague priors for model selection can be produce very misleading results, and will typically have the unintended consequence of sending the Bayes factor in favor of the simpler model to infinity. This is often known as Bartlett’s paradox in the simple context of testing normal means (Bartlett, 1957), but the same principle applies here.

It may also be the case that elements of $\Omega$ depend upon some parameter $\eta$. Since this parameter appears only in the alternative model, $\eta$ needs a proper prior, or else the marginal likelihoods will be defined only up to an arbitrary multiplicative constant.

Similar challenges occur in all model-selection problems. General approaches and guidelines for choosing priors on nonshared parameters can be found in Laud and Ibrahim (1995), (O’Hagan, 1995), (Berger and Pericchi, 1996), and (Berger et al., 1998). But very few tools of analogous generality have been developed for nonparametric problems, with most work concentrating on how to compute Bayes factors for pre-specified models (Basu and Chib, 2003), or how to test a parametric null against a nonparametric alternative of a suitably restricted form (Berger and Guglielmi, 2001).

This leaves just two obvious criteria for choosing $\Omega$ and $F$ in the face of weak prior information:

1. Elements of $\Omega$ should be smooth, i.e. slowly varying on the unit-time scale of the residual model. This will allow deconvolution of the mean process from the residual, and reflects the prior belief that the
mean function will describe long-term departures from 0 in the face of short-term autoregressive jitters. (Indeed, these departures are precisely what the methodology is meant to detect.)

2. $F$ should be centered at the null model, and should concentrate most of its mass on elements of $\Omega$ that predict $y$ values on a scale similar to those predicted by the null model. This will avoid Bartlett’s paradox, and generalizes the argument made by Jeffreys (1961) in recommending an appropriately scaled Cauchy prior for testing normal means.

These criteria allow much wiggle room, but at least provide a starting point. Unfortunately there is no objective solution, in this or in any model-selection problem, though the closest thing to a default approach is to simply choose the marginal variance of the alternative process to exactly match the marginal variance of the null process. Best, of course, is to conduct a robustness study, where the features of the nonparametric alternative not shared by the null are varied in order to assess changes in the conclusions. This will usually be quite difficult in large multiple-testing problems, since computations for just a single version of the alternative model may be expensive.

The choice of $\alpha$, the precision parameter for the residual Dirichlet-process prior, is relatively free by comparison, since this parameter appears in both the null and alternative models. Strictly speaking, in order to use a noninformative prior for $\alpha$, verification of the conditions in Berger et al. (1998) regarding group invariance is necessary, which is difficult in this case. (The issue is that a parameter does not necessarily mean the same thing in both $M_0$ and $M_A$ just because it is assigned the same symbol in each.) In the absence of a formal justification for using a noninformative prior, the conservative approach is to elicit priors for $\alpha$ in terms of the expected number of AR(1) mixture components in each of $M_0$ and $M_A$. Often there will be extrinsic justification for choosing $\alpha$ to be the same under both models.

5. A model for the corporate-performance data.

5.1. Model details. As an example of how tests involving (11)–(13) can be constructed, this section outlines a nonparametric model for a larger subset of the corporate-performance data containing 5,498 firms. This contains every publicly traded American company between 1965 and 2005 for which at least 15 years of history were available.

The class of Gaussian processes with some known covariance function is ideally suited for modeling nonzero trajectories, since the covariance function can be chosen to yield smooth functions with probability 1, and since the prior marginal variance of the process can be controlled exactly (so
that Bartlett’s paradox may be easily avoided). Gaussian processes have the added advantage of analytical tractability, which is very important in hypothesis testing because of the need to evaluate the marginal likelihood of the data under the alternative model. More general classes of functions are certainly possible, though perhaps computationally challenging in the face of massive multiplicity.

One additional feature to account for is clustering, since management theorists are interested in identifying a small collection of archetypal trajectories that may correspond to different sources of competitive advantage. Partitioning of firms into shared trajectories is especially relevant for advocates of the so-called “resource-based view” of the firm (Wernerfelt, 1984). Additionally, clustering on treatment effects is known to increase power in multiple-testing problems (Dahl and Newton, 2007).

The approach considered here is similar to that introduced by Dunson and Herring (2006). The distribution of nonzero random functions is modeled with a functional Dirichlet process:

\( F \sim \text{FDP}(\nu, \text{GP}(C_\kappa)) \) (14)

\( C_\kappa(t_1, t_2) = \kappa_1 \cdot \exp\left(-0.5 \cdot \frac{|t_1 - t_2|}{\kappa_2}\right)^2. \) (15)

The functional Dirichlet process in (14) has precision parameter \( \nu \) and is centered at a Gaussian process with covariance function \( C_\kappa \). At time \( t \), the value of the function \( f_i \) has a Dirichlet-process marginal distribution: \( f_i(t) \sim \text{DP}(\nu, N(0, \kappa_1)) \). In choosing the hyperparameter \( \kappa \), close attention must be paid to the marginal variance of the residual model, so that variance inflation in (15) does not overwhelm the Bayes factor. For greater detail on Gaussian processes for nonparametric function estimation, see Rasmussen and Williams (2006).

This model is significantly richer than the simplistic framework developed in Section 2, but is similar in two crucial ways:

**Centering at the null model,** since the Gaussian process in (15) leads to \( \text{E}(f_i | \gamma_i = 1) = 0 \). As before, it is equally likely \textit{a priori} that a firm’s trajectory will be predominantly negative or predominantly positive.

**Variance inflation** under the alternative model is controlled through the choice of a single hyperparameter, with \( \kappa_1 \) in (15) playing the role of \( \sigma^2 \) in (3). Hence despite the complicated nonparametric wrapper, the Ockham’s-razor effect upon the implied marginal likelihoods still happens in the familiar way.

The model also solves both of the major problems encountered in the ROA
data: time-varying nonzero trajectories, and clustering both of trajectories and of company-specific parameters for the autoregressive residual.

An extensive prior-elicitation process was undertaken with three experts in management theory who had originally compiled the data. For the base measure of the Dirichlet-process mixture of AR(1) covariance models, the same hyperparameters from the parametric model in (6) and (7) were used. Hence the starting point for this elicitation was: $\kappa_1 \approx 1.94$, which is the prior marginal variance of the residual AR process (assessed by simulation), and $\kappa_2 = 15$ (on a 41-year time scale), which reflected the experts’ judgments about the long-term effects of strategic choices made by firms. The elicitees were repeatedly shown trajectories drawn from this prior and other similar priors, and soon settled upon $\kappa_1 = 1.25$ and $\kappa_2 = 13$ as values that better reflected their expectations. Additionally, they chose $\alpha = 10/\log N$, and $\nu = 15/\log N$ on the basis of how many clusters they expected.

For the actual data, a third trajectory model was also introduced: a DP mixture of constant trajectories rather than of Gaussian-process trajectories. This entails only a slight complication of the analysis, in that now (13) is a three-component mixture rather than a two-component mixture. This is equivalent to including the limiting-linear-model framework of Gramacy (2005)—whereby a flat trajectory is given nonzero probability as an explicit limiting case of the Gaussian process—inside the base measure of the functional Dirichlet process itself.

5.2. Results. The above model was implemented using the blocked Gibbs-sampling algorithm of Ishwaran and James (2001) to draw from the nonparametric distributions $F$ and $G$. Convergence was assessed through multiple restarts from different starting points, and was judged to be satisfactory.

Overall, 981 of 5,498 firms were flagged as being from the alternative model with greater than 50% probability, representing an overall discovery rate of about 18%. Of these, only 196 firms were from the alternative model with greater than 90% probability.

To assess robustness to the hyperparameter choices $\kappa_1$ and $\kappa_2$, which control the marginal variance and temporal range of of the Gaussian process base measure in (15), the results were recomputed for a coarse grid of 12 pairs of values spanning $0.75 \leq \kappa_1 \leq 2.25$ and $5 \leq \kappa_2 \leq 20$. This reflected the lower and upper ends of what the collaborating management theorists considered reasonable on the basis of observing draws from these priors.

As expected, larger values of $\kappa_1$ tended to yield fewer non-null classifications (due to variance inflation in the marginal likelihoods), while larger values of $\kappa_2$ tended to punish firms whose peaks and valleys in performance

Nonparametric Multiple Testing of Time Series

Table 2

Posterior probabilities for six firms that were flagged as being from trajectory GP 8 with greater than 50% probability in the MLE clustering analysis; see Figure 3 for labels. (GV refers to a unique corporate identifier.)

<table>
<thead>
<tr>
<th>GV</th>
<th>Company Name</th>
<th>Flat 1</th>
<th>Flat 2</th>
<th>GP 7</th>
<th>GP 8</th>
<th>Other</th>
<th>Null</th>
</tr>
</thead>
<tbody>
<tr>
<td>11535</td>
<td>Winn-Dixie Stores</td>
<td>4</td>
<td>1</td>
<td>10</td>
<td>78</td>
<td>5</td>
<td>2</td>
</tr>
<tr>
<td>4828</td>
<td>Delhaize America</td>
<td>7</td>
<td>2</td>
<td>11</td>
<td>75</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>6830</td>
<td>Lubrizol Corp</td>
<td>20</td>
<td>9</td>
<td>2</td>
<td>64</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>7139</td>
<td>Maytag Corp</td>
<td>25</td>
<td>2</td>
<td>8</td>
<td>57</td>
<td>2</td>
<td>6</td>
</tr>
<tr>
<td>4323</td>
<td>Emery Air Freight</td>
<td>5</td>
<td>2</td>
<td>29</td>
<td>57</td>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>7734</td>
<td>National Gas &amp; Oil</td>
<td>18</td>
<td>13</td>
<td>11</td>
<td>53</td>
<td>3</td>
<td>2</td>
</tr>
</tbody>
</table>

were short-lived over the 41-year time horizon. Many firms that were borderline in the original analysis (that is, having $p_i$ just barely larger than 50%) were reclassified as “noise” for certain other values of $\kappa$. Yet a stable cohort of 246 firms were flagged as non-null in all 12 analyses, suggesting a reasonable degree of robustness with respect to hyperparameter choice.

The behavior of individual firms was then characterized by using the MCMC history from the original analysis to get a maximum-likelihood estimate of the nonparametric alternative model. The almost-sure discreteness of the Dirichlet process means that this estimate is a mixture of a small number of flat and Gaussian-process trajectories. The 17 highest-weight trajectories in the MLE are in Figure 3; they are split into four loose categories reflecting different archetypes of company performance.

This allows an MLE clustering analysis: if $F$ in (14) is frozen at the mixture model in Figure 3 and the MCMC rerun, it is possible to flag companies that come from specific clusters. (Strictly speaking, only the first 17 atoms in the stick-breaking approximation of $F$ were frozen; others atoms were still considered, but they were allowed to vary.) Examples of the kinds of summaries available are in Table 2 and Figure 4.

Some general features of the methodology are apparent from these results:

- There is substantial shrinkage of estimated mean trajectories back toward the global average (i.e. the 50th percentile). This is itself a form of multiplicity correction, in that extreme outcomes are quite likely to be attributed to chance even among those firms flagged as being from the alternative model.
- Often there is no dominant trajectory in the MLE cluster set that characterizes a specific firm’s history. For the three firms in Figure 4, the probability is split among two or even three trajectories.
Fig 3. The 17 highest-weight trajectories in the MLE estimate of the alternative model, split into four loose categories. The y-axis is given on the $N(0,1)$ inverse-cdf scale to reflect the quantile of each firm’s performance.
Fig 4. Maytag, Plenum Publishing, and El Paso CGP: actual ROA histories along with trajectory-membership probabilities from the MLE clustering analysis.
Model-averaged predictions of future performance are available with very little extra work, since full MCMC histories of each firm’s trajectory and residual model are available.

The MLE clustering analysis can provide evidence for historical evolution within specific firms, which is of great interest as the subject of follow-up case studies. Maytag, for example, displays markedly different performance patterns before and after 1987, which is reflected in its high probability of being from a falling trajectory (GP 8).

It must be emphasized that any such clustering analysis is at best an approximation, aside from the fact that the models themselves are also approximations. It relies, after all, upon a single point estimate of the trajectories composing the random distribution \( F \), and as such ignores uncertainty about the trajectories themselves. In the example at hand, several independent runs were conducted; each yielded a different MLE cluster set, but the same broad patterns (e.g. something like GP 8, something like Flat 3, and so on) emerged each time, suggesting at least some degree of robustness of the qualitative conclusions.

Still, the most reliable quantities are the inclusion probabilities \( \{p_i\} \) computed from the full nonparametric analysis, which should form the basis of claims regarding which units are from the null and which are not.

6. Type-I error performance: a simulation study. This section recapitulates the simulation study of Section 2.3 using the more complicated models. The goal is to assess Type-I error performance by applying the methodology outlined here to a simulated data set where the number of nonzero trajectories is known.

The true residual model \( G \) was constructed using a single MCMC draw from the stick-breaking representation of the nonparametric residual model for the corporate-performance data. This corresponded roughly to a 21-component mixture of AR(1) models (since 39 of the 60 atoms in the stick-breaking representation of \( G \) had trivial weights), with many \((\phi,v)\) pairs that differed starkly in character.

To simulate the true mean trajectories, two independent sets of 5,500 draws were taken from the prior in (11)-(13). In the first simulated set of trajectories, the true value of \( p \) was fixed at 1/5 in order to roughly approximate the fraction of discoveries on the ROA data. In the second set, \( p \) was fixed at 1/55, to reflect a much sparser collection of signals. Upon sampling, these yielded 1126 and 98 nonzero trajectories, respectively.

In both studies, each of these trajectories was convolved with a single independent vector of autoregressive noise drawn from the true \( G \)—in other
words, with a highly complex pattern of residual variation of the type that was shown to flummox the “single AR(1)” model of Section 2. For the sake of comparison, the same set of 5500 noise vectors was used for each experiment.

The results of these simulations were very encouraging for the Type-I error performance of the more complicated model. In the first simulated data set, 297 of the 1126 nonzero trajectories were flagged as non-nulls with greater than 50% probability. Only 24 of the 4374 null companies were falsely flagged, and of these, only 3 had larger than a 70% inclusion probability.

In the second simulated data set, of the 98 known nonzero trajectories, 24 had posterior inclusion probabilities greater than 50%, and 6 had inclusion probabilities greater than 90%. Of the 5402 null trajectories, only 2 had inclusion probabilities greater than 50% (these two were only 63% and 67%).

If $p_i \geq 0.5$ is taken as the decision rule for a posterior classification of a trajectory as non-null (which again reflects a symmetric 0–1 loss function), then the realized false-discovery rates were only 7.7% on 26 discoveries in the sparser case, and only 7.5% on 321 discoveries in the denser case. (The closeness of these two realized false-discovery rates may simply be a coincidence of the particular values of $p$ chosen).

This suggests that a sizeable fraction of the 981 firms flagged as non-null trajectories in Section 5.2 represent non-average performers, and are not false positives.

7. Summary. This paper has described a framework for Bayesian multiple hypothesis testing in time-series analysis. The proposed methodology requires specifying only a few key hyperparameters for the nonparametric null and alternative models, and general guidelines for choosing these quantities have been given. Importantly, no ad-hoc “correction factors” are necessary in order to introduce a penalty for multiple testing. Rather, this penalty happens quite naturally by treating the prior inclusion probability $p$ as an unknown quantity with a noninformative prior.

Naive characterizations of the null hypothesis are shown to have poor error performance, suggesting that the Bayesian procedure is highly sensitive to the accuracy of the null model used to describe an “average” time series. Yet once a sufficiently complex model for residual variation is specified, the procedure exhibits very strong control over the number of false-positive declarations, even in the face of firms with different autoregressive profiles. The difference between the results in Section 2.3 and Section 6 highlights the effectiveness and practicality of using nonparametric methods as a general error-robustification tactic in multiple-testing problems.

Posterior inference for a specific time series can be summarized in at least
three ways: by quoting $p_i$ (the probability of that unit’s being from the alternative model), by performing an MLE clustering analysis as in Section 5.2, or by plotting the posterior draws of the unit’s nonzero mean trajectory. Plots such as Figure 4 can be quite useful for communicating inferences to nonexperts, as in the management-theory example considered throughout.

The companies flagged as impressive performers, of course, can only be judged so with respect to a particular notion of randomness: the DP mixture of autoregressive models described in Section 3. Inference on the nonzero trajectories can still reflect model misspecification, and cannot unambiguously identify companies that have found a source of sustained competitive advantage. This Dirichlet-process model, however, is a much more general statement of the null models postulated by Denrell (2003) and Denrell (2005), suggesting that the procedure described here can identify firms that, with high posterior probability, depart from randomness in a specific way that may be interesting to researchers in strategic management.

References.


