Sequential algorithmic modification with test data reuse

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Abstract

After initial release of a machine learning algorithm, the model can be fine-tuned by retraining on subsequently gathered data, adding newly discovered features, or more. Each modification introduces a risk of deteriorating performance and must be validated on a test dataset. It may not always be practical to assemble a new dataset for testing each modification, especially when most modifications are minor or are implemented in rapid succession. Recent works have shown how one can repeatedly test modifications on the same dataset and protect against overfitting by (i) discretizing test results along a grid and (ii) applying a Bonferroni correction to adjust for the total number of modifications considered by an adaptive developer. However, the standard Bonferroni correction is overly conservative when most modifications are beneficial and/or highly correlated. This work investigates more powerful approaches using alpha-recycling and sequentially-rejective graphical procedures (SRGPs). We introduce novel extensions that account for correlation between adaptively chosen algorithmic modifications. In empirical analyses, the SRGPs control the error rate of approving unacceptable modifications and approve a substantially higher number of beneficial modifications than previous approaches.

1 INTRODUCTION

Before a machine learning (ML) algorithm is approved for deployment, its performance is usually evaluated on an independent test dataset. If the ML algorithm is modified over time, its performance may change. There are no guarantees on how the performance may evolve when the model developer is allowed to introduce modifications in an unconstrained manner. For instance, algorithmic modifications that reduce computational costs may negatively impact model accuracy or precision, and improvements along an aggregate performance metric may come at the cost of worse performance for certain minority subgroups and exacerbate issues of algorithmic fairness. To check that a proposed modification is acceptable for deployment, the current approach is to run a hypothesis test on a new test dataset, separate from the original one (Feng et al., 2020). The null hypothesis is that the modification is not acceptable; a modification is approved if we successfully reject the null. Nevertheless, large high-quality test datasets are often hard to acquire, particularly in the medical setting. A major motivation for this work comes from the FDA’s recent interest in letting medical device developers update ML-based software, while still ensuring its safety and effectiveness (U.S. FDA, 2019).

When labeled data are expensive and/or difficult to collect, it is tempting to reuse an existing test dataset for determining the acceptability of an algorithmic modification. The danger of test data reuse is that the model developer can learn aspects of the test data when it is used in a sequential and adaptive manner, creating dependencies between algorithmic modifications and the holdout data. For instance, the model developer may inadvertently incorporate spurious correlations in the test data to attain over-optimistic performance estimates. This feedback loop introduces bias to the performance evaluation procedure, and adaptively defined hypothesis tests can have drastically inflated Type I error rates (Gelman and Loken, 2017; Thompson et al., 2020).

Recent works protect against inappropriate test data reuse and overfitting by reducing the amount of information released by the testing procedure (Russo and Zou, 2016). The two main approaches are to either coarsen the test outputs along a grid of values (Blum and Hardt, 2015; Rogers et al., 2019) or to perturb the test results with random noise using differential privacy techniques (Dwork et al., 2015a; Feldman and Steinke, 2018). However, existing methods require immensely large datasets to provide protection against overfitting with theoretical guarantees (Rogers et al., 2019).

Our aim is to design valid test data reuse procedures for smaller sample sizes that still have sufficiently high power.
to approve good algorithmic modifications. Our focus is on methods that coarsen the test results. In fact, we consider the extreme case of coarsening where the procedure releases a single bit of information, e.g. whether or not the modification was approved.

When test results are coarsened, the adaptive modification strategy can be described as a tree. As such, one can view the test data reuse problem as a multiple hypothesis testing problem: If we control the family-wise error rate across the entire tree, we control the probability of approving one or more unacceptable modifications. Existing procedures perform a Bonferroni correction with respect to the size of this tree (Blum and Hardt [2015], Rogers et al. [2019]). Nevertheless, the Bonferroni correction is known to be conservative. Instead, we can gain significant power using alpha-recycling (Burman et al. [2009]) and accounting for correlation between test statistics (Westfall and Stanley Young [1993]). Indeed, we expect algorithmic modifications to be highly correlated when there is significant overlap between their training data and similarities in their training procedures.

In this paper, we design valid test data reuse procedures based on sequentially rejective graphical procedures (SRGPs) (Bretz et al. [2009, 2011a,b]). Although SRGPs are well-established technique for testing many pre-specified hypotheses, many of these procedures cannot be applied when the hypotheses are adaptively defined in sequence. The main challenge is that many nodes in the tree of hypotheses are not observed. As such, we introduce two novel SRGPs that are able to account for correlation between adaptively-defined algorithmic modifications without needing to observe these “counterfactual” hypotheses. The first procedure accounts for correlation between observed nodes in the tree using a fixed-sequence testing procedure. The second procedure is based on the fact that analysts are not adversarial in practice, i.e. they will not purposefully use prior results to overfit to the test data (Mania et al. [2019], Zrnic and Hardt [2019]). We leverage this fact by requiring the model developer to pre-specify a hypothetical online learning procedure. The SRGP then utilizes the similarity between the adaptive and pre-specified modifications to improve testing power. Both methods can be applied to test black-box ML algorithms and, importantly, are suitable for smaller sample sizes, e.g. those commonly found in medical settings. In empirical analyses, our procedures protect against overfitting to the test data and approve a higher proportion of acceptable modifications than existing approaches. Code will be publicly available on Github.

2 PROBLEM SETUP

Suppose the test dataset is composed of $n$ independently and identically distributed (IID) observations $(X_1, Y_1), (X_2, Y_2), \ldots, (X_n, Y_n) \in X \times Y$ drawn from the target population. Consider a model developer who adaptively proposes a sequence of $T$ algorithmic modifications \{\hat{f}_1^{\text{adapt}}, \ldots, \hat{f}_T^{\text{adapt}}\}, where each modification is a model that predicts some value in $Y$ given input $X$. Given criteria for defining the acceptability of a modification, our goal is to approve as many acceptable modifications as possible while controlling the probability of approving an unacceptable modification. Because the decision to approve a modification can be framed as a hypothesis test, a procedure for approving adaptively-defined modifications is equivalent to testing a sequence of adaptively-defined hypotheses $H_1^{\text{adapt}}, \ldots, H_T^{\text{adapt}}$. Moreover, control of the online family-wise error rate (FWER) in the strong sense, i.e. for any configuration of the null hypotheses, implies control over of the rate of approving at least one unacceptable modification.

There are various ways to define acceptability and their corresponding hypothesis test. For example, we may define a modification $f$ to be acceptable as long as its expected loss is smaller than that of the original model $f_0$. So given a real-valued loss function $\ell$, we would test the hypothesis

$$H_j^{\text{adapt}} : \mathbb{E} \left( \ell \left( f_j^{\text{adapt}}(X), Y \right) \right) \geq \mathbb{E} \left( \ell \left( f_0(X), Y \right) \right)$$

at each iteration $j = 1, \ldots, T$. If we require monotonic improvement in the model performance, we can test if the $j$-th modification is superior to the most recently approved modification $f_j^{\text{approved}}(X)$, i.e.

$$H_0^{\text{adapt}} : \mathbb{E} \left( \ell \left( f_j^{\text{adapt}}(X), Y \right) \right) \geq \mathbb{E} \left( \ell \left( f_j^{\text{approved}}(X), Y \right) \right).$$

Finally, one may also consider multidimensional characterizations of model performance (e.g. model performance within subgroups) and define acceptability as a combination of superiority and non-inferiority tests (Feng et al. [2020]). The testing procedures described below only depend on the $p$-values, so we leave the specific definition of acceptability unspecified until the experimental section.

To limit the adaptivity of the model developer, we consider procedures that sequentially release a single bit of information for each test: One means that the modification is approved and zero means it is not. Consequently, modifications proposed by any adaptive strategy can be described as a bifurcating tree, where $f_0$, is the nonadaptive model tested at time $t$ for the history of approvals $a_t \in \{0, 1\}^{t-1}$, $H_{a_t}$ is the associated nonadaptive hypothesis test, and $p_{a_t}$ is its marginal $p$-value. While one can regard this set of $(2^T - 1)$ hypotheses tests as prespecified, we are only able to observe a specific path along this tree. The unobserved hypotheses are counterfactuals. As such, we need a multiple testing procedure (MTP) that controls the FWER for any adaptively chosen path along the tree without knowing the exact nature of the counterfactual hypothesis tests.

A simple approach is to perform a uniform Bonferroni correction for the size of the entire tree. However, the standard Bonferroni procedure has low power because it ignores correlations between models and allocates substantial test mass
We can design more powerful test data reuse procedures by building on sequentially rejective graphical procedures (SRGPs), which use directed graphs to define a wide variety of iterative MTPs such as gatekeeping procedures, fixed sequence tests, and fallback procedures [Bretz et al., 2009]. SRGPs traditionally assume the set of hypotheses \{H_j : j \in I\} is prespecified and known. The graph initially contains one node for each elementary hypothesis \(H_j\), where each node is associated with a non-negative weight \(w_j(I)\). The initial node weights, which are constrained to sum to one, control how the total alpha is divided across the elementary hypotheses and correspond to a set of adjusted significance thresholds \(c_j(I)\). We reject elementary hypothesis \(H_j\) in the current graph if its marginal p-value \(p_j\) is smaller than \(c_j(I)\).

For instance, a standard Bonferroni correction is represented by the initial weights of \(w_j(I) = 1/|I|\) for all \(j \in I\) and significance thresholds \(c_j(I) = w_j(I)\alpha\). In addition, the graph contains directed edges where the edge \(H_j\) to \(H_k\) is associated with weight \(g_{j,k}(I)\) for \(j, k \in I\) and edge weights starting from the same node must sum to one. When an elementary hypothesis \(H_j\) is rejected, its node is removed from the graph and its weight is propagated to its children nodes. This redistribution of test mass, also known as alpha-recycling, increases the power for testing the remaining hypotheses and strictly improves upon simpler procedures that do not use recycling. More specifically, the weight of the edge from \(H_j\) to \(H_k\), denoted \(g_{j,k}(I)\), represents how much of \(H_j\)’s node weight will be redistributed to \(H_k\) if \(H_j\) is rejected. So when \(H_j\) is removed, the new weight for hypothesis \(H_k\) for \(k \in I \setminus \{j\}\) is

\[
    w_k(I') = w_k(I) + g_{j,k}(I)w_j(I).
\]

Outgoing edges for all remaining nodes are also renormalized to sum back to one. The SRGP continues until no more hypotheses can be rejected. See Figure 1 for an example.

Assuming all test reports are binary, we can describe adaptive test data reuse as the following prespecified SRGP. Let \(I_t\) be the set of hypotheses remaining at time \(t\), where \(I_0\) is the initial set. We only consider SRGPs with nonzero edge weights from hypothesis \(H_{a_t}\) to the sequence of hypotheses that would be tested upon rejection of \(H_{a_t}\) but prior to the next rejection (though one may also consider more complex recycling procedures). For such SRGPs, the graph of hypotheses has the tree structure seen in Figure 2a, where the only edges in the tree are between hypotheses \(H_{a_t}\) and \(H_{a_{t'}}\) for \(t < t'\) and

\[
    a_{t',j} = a_{t,j}1\{j < t\} + 1\{j = t\}.
\]

Note that the SRGP tree is not the same as the bifurcating tree for generating hypotheses, as the former describes how alpha is recycled.

The model developer must prespecify all initial node weights \(w_{a_t}(I_0)\). A simple approach is to perform a uniform Bonferroni correction across all nodes in the graph. We can achieve more power by assigning larger weights to nodes that are more likely to be tested. For example, if the model developer knows that all their modifications will be approved, they should set the initial node weight for \(H_1\) and all edge weights along the top path in Figure 2a to one.

**Algorithm 1** A sequentially rejective graphical procedure (SRGP) that only outputs binary test reports for \(T\) adaptive hypotheses given function \(\text{compute sig threshold}\).

**Require:** Initialize \(I_0\) as the set of all nodes in the prespecified tree; initialize \(a_1 = ()\) and \(\tau_1 = 0\); choose node weights \(w_{a_t}(I_0)\) for all \(t = 1, 2, \ldots, T\) and \(a_t' \in \{0, 1\}^{t-1}\); and set \(w_{a_0}(I_0) = 0\).

**Ensure:** \(\sum_{a_t'} w_{a_t'}(I_0) = 1\).

**for** \(t = 1, 2, \ldots, T\) **do**

Specify edge weight \(g_{a_{t},a_t}\) that satisfies outgoing edge weight constraints.

**Weight propagation**

\[
    w_{a_t}(I_t) = w_{a_t}(I_{t-1}) + g_{a_{t},a_t}w_{a_{t'}}(I_{\tau_t})
\]

for all \(t', a_{t'}\) such that \(H_{a_{t'}} \in I_t\) and \(a_t' \neq a_t\) do

**# Other weights remain unchanged**

\[
    w_{a_{t'}}(I_t) = w_{a_{t'}}(I_{t-1})
\]

end for

Let \(p_{a_t}\) be the marginal p-value from testing \(H_{a_t}\). Compute significance threshold \(c_{a_t}(I_t)\) using \(\text{compute sig threshold}(a_t, \{w_{a_t}(I_t) : t', a_{t'}\})\) if \(p_{a_t} \leq c_{a_t}(I_t)\) then

Report that \(j_{t,\text{adapt}}\) has been approved.

**# Remove node**

\[
    I_{t+1} = I_t \setminus a_t
\]

\[
    \tau_{t+1} = t
\]

\[
    a_{t+1} = (a_t, 1)
\]

else

Report that \(j_{t,\text{adapt}}\) has not been approved.

\[
    \tau_{t+1} = \tau_t
\]

\[
    a_{t+1} = (a_t, 0)
\]

end if

end for

The model developer will only need to incrementally re-
we can perform the closed test using a sequentially rejective weight can be treated as a constant because the only relevant edge weight at time \( t \) is \( g_{a_t,a_t}(I_t) \) and its value is equal to \( g_{a_t,a_t}(I_{t'}) \) for all \( t' < t \). As such, this procedure for specifying node and edge weights corresponds to a fully prespecified SRGP where a subset of the edge weights are revealed sequentially. To make sure that this SRGP can be executed in the adaptive setting, we must be able to calculate the adjusted significance thresholds for the adaptive hypotheses given the current set of node weights without observing the counterfactual hypotheses.

The entire SRGP algorithm for testing adaptive algorithmic modifications is outlined in Algorithm 1. It accepts some function compute\_sig\_threshold that outputs the significance threshold for the adaptively chosen hypothesis given node weights in the current tree. To prove that an SRGP with function compute\_sig\_threshold controls the FWER, we must show that it is a closed test procedure that satisfies the consonance property. Recall that a closed test procedure uses the following recipe to control the FWER at level \( \alpha \): it rejects an elementary hypothesis \( H_j \) if the intersection hypothesis \( H_K = \cap_{k \in K} H_k \) for every subset \( K \subseteq I \) containing the elementary hypothesis \( H_j \) is rejected at level \( \alpha \) (Lehmann and Romano, 2005). Moreover, a closed test satisfies the consonance property if the following is true for all \( J \subseteq I \): if intersection hypothesis \( H_J \) is rejected locally (i.e., its \( p \)-value is no more than \( \alpha \)), there exists some \( j \in J \) such that \( H_K \) can be rejected locally for all \( K \subseteq J \) with \( j \in K \) (Gabriel, 1969). In particular, it follows that the corresponding elementary hypothesis \( H_j \) can be rejected by the closed test procedure. When consonance holds, we can perform the closed test using a sequentially rejective (or “shortcut”) procedure that iteratively rejects the elementary hypotheses without needing to test every intersection hypothesis (Hommel et al., 2007). When hypothesis tests are fully prespecified, consonance makes closed testing more computationally efficient/tractable. The consonance property is even more important in the adaptive setting because we can reject the adaptive hypotheses without observing counterfactual or future hypotheses. As such, the consonance property of an SRGP in the adaptive setting is not simply for computational efficiency, but is necessary for being able to compute anything.

Below, we will describe three SRGPs for testing an adaptive sequence of algorithmic modifications, presented in order of increasing complexity. Each differ in how compute\_sig\_threshold is defined. To prove that the procedures satisfy the consonance property, it is sufficient to show that the following monotonicity condition holds (Bretz et al., 2009): For every pair of subsets \( K, J \subseteq I \) where \( K \subseteq J \) and \( j \in K \), we have

\[
c_j(J) \leq c_j(K). \tag{3}
\]

All proofs are provided in the Appendix.

### 2.2 BONFERRONI-BASED SRGPS

We begin with the simplest SRGP that performs closed testing with a weighted Bonferroni-Holm correction based on node weights, which was originally proposed in Bretz et al. (2009) to test a set of fully pre-specified hypotheses. Nevertheless, this procedure can also be applied in the adaptive setting because the significance thresholds do not depend on observing the counterfactual hypotheses. In particular, this procedure tests the \( t \)-th adaptive hypothesis given history \( a_t \) by comparing its marginal \( p \)-value to the corrected significance threshold \( c_{a_t}(I_t) = w_{a_t}(I_t) \alpha \). Because this closed test satisfies the monotonicity condition, Algorithm 1 with this significance threshold controls the FWER for the adaptive hypotheses at level \( \alpha \).

As a simple example, consider an SRGP that initially assigns Bonferroni-corrected weights to every node and selects nonzero edge weights. This is more powerful than per-
forming a standard Bonferroni correction without any alpha-recycling because the significance thresholds are monotonically non-decreasing at each iteration.

2.3 SRGPs with Fixed Sequence Tests for Correlated Modifications

In practice, algorithmic modifications are likely to be highly correlated. In this case, a Bonferroni-based SRGP is conservative. We can design more powerful SRGPs by taking into account correlation between the p-values. Bretz et al. (2011a) proposed a procedure that calculates an inflation factor $c(I)$ for intersection hypothesis $I$ such that the probability there exists an elementary hypothesis $H_{ij}$ with marginal p-value $p_{ij}$ less than $c(I)w_{ij}(I)\alpha$, under the null $I$, is no more than $\alpha$. Millen and Dmitrienko (2011) proposed a similar procedure but for test statistics and critical values. Unfortunately, both procedures require knowing the exact correlation structure between all the hypotheses and checking that the monotonicity property holds. This is not feasible in the adaptive setting. To resolve these issues, we propose a new SRGP that (1) partitions the hypothesis tree into sequences of observed hypotheses and (2) uses a fixed-sequence test within each subgroup.

We group together hypotheses that would be tested along a streak of failures immediately following a successful approval (Figure 2B). That is, we define a subgroup for history $a_t \in \{0, 1\}^T$ with $a_{t-1} = 1$ as the hypotheses with histories $a_{t'} = (a_t, \vec{0})$ for any length zero vector, i.e.

$$G_{a_t} = \{ H_{a_{t'}}, a_{t',i} = a_{t,i} \, \forall \, i \in \{1, ..., t'-1\}, \forall t' \geq t \}$$

To test intersection hypothesis $I$, we test each subgroup $G_{a_t} \cap I$ at level \(\sum_{H_{a_{t'}}, a_{t',i} = a_{t,i}} w_{a_{t'},I} \alpha\). We reject $H_I$ at level $\alpha$ if any of the subgroup-specific tests are rejected. We can show that this controls the Type I error at level $\alpha$ using a union bound. To test a subgroup, we test its hypotheses in the order they are revealed and spend up to the allocated alpha weight. To satisfy the monotonicity property, the significance threshold $c_{a_t}(I)$ for $a_t \in G_{a_t} \cap I$ is defined as the maximum threshold that spends no more than the allocated alpha up to time $j$ for all subsets of hypotheses, i.e.

$$c_{a_t}(I) = \sup \{\tilde{c} \mid \Pr(p_{a_k} > c_{a_k}(I) \forall k \in K, p_{a_k} < \tilde{c} | H_{K \cup \{a_t\}}) \leq \sum_{a_k \in (G_{a_{t'}} \cap I \setminus K) \setminus K, k \leq j} w_{a_{t'},I} \alpha, \forall K \subseteq \{a_k : a_k \in G_{a_t} \cap I, k < j\} \}$$

This expression is complicated because it handles arbitrary correlation structures between the p-values. It greatly simplifies in certain cases. For example, if we are performing one-sided Z-tests and the pairwise correlations of the model losses are non-negative, (4) is equivalent to defining $c_{a_j}(I)$ as the solution to

$$\Pr(p_{a_k} > c_{a_k}(I) \forall k = t, ..., j - 1, p_{a_j} < c_{a_j}(I) | \cap_{k=t}^{j-1} H_{a_k}) = w_{a_j}(I)\alpha.$$ 

Using the fixed sequence tests from above, we sequentially calculate the significance thresholds and test the adaptive hypotheses. When a hypothesis is rejected, we remove its node and propagate its local weight to its children nodes per (I). We can prove the monotonicity condition holds to establish the following result:

**Theorem 1.** Algorithm [I] with significance thresholds chosen using (4) controls the FWER for adaptively defined hypotheses at level $\alpha$.

2.4 SRGPs with Prespecified Hypothetical Model Updates

The SRGPs in the above sections protect against the worst case scenario where the model developer is adversarial. In practice, the model developer may have a plan for how they will update their model over time (i.e. continually refit the model on accumulating data) and will only make small adjustments based on test results. As such, we do not expect the adaptively chosen model at iteration $t$ to stray far from the initial plan. In the most extreme case, we may find that the model developer is not adaptive at all and follows the prespecified procedure perfectly; instead of correcting for $(2^T - 1)$ hypotheses, we would expect that the correction factor to be $O(T)$ instead.

To leverage this similarity assumption, we propose a novel SRGP that requires the model developer to prespecify a procedure for generating hypothetical model updates. This prespecified procedure describes the exact steps for how modifications would be generated, e.g. the data stream used, the number of training observations, and hyperparameter selection. These hypothetical model updates are included as additional nodes in the hypothesis graph and assigned positive node weights. Their sole purpose is to improve power for approving the adaptively-defined model updates. These model updates are never formally tested nor approved for deployment. We also do not release any information about their test performance, because doing so would increase the amount of information leaked to the model developer and the branching factor of the adaptive tree.

At each iteration, this SRGP constructs a confidence region for the performance of the $t$-th prespecified model update $f^{\text{pres}}_{t}$ by spending its allocated alpha, accounting for its correlation with all prespecified models up to iteration $t - 1$. It then tests the $t$-th adaptive model by accounting for its correlation with the prespecified models up to iteration $t$. As such, the power for testing the adaptive modifications increase as their correlation with the prespecified updates increases.
More formally, the critical value and significance threshold at time $t$ are calculated as follows. Let $P_0$ denote the target population and $P_n$ denote the empirical distribution of the test dataset. Here we consider a univariate performance measure $\psi$, where $\psi\left(\hat{f}, P\right)$ is the performance of model $\hat{f}$ with respect to distribution $P$. It is straightforward to extend this procedure to multivariate performance measures (see the Appendix for an example). Denote the deviation between the estimated and true performance as

$$\xi_{t,n}^\text{pres} = \psi\left(\hat{f}_t^\text{pres}, P_n\right) - \psi\left(\hat{f}_0^\text{pres}, P_0\right).$$

For intersection hypothesis $I$, define $\bar{I}$ as union of $I$ and all prespecified nodes. Define critical value $z_{t}^\text{pres}(I)$ as the largest $\tilde{c}$ such that

$$\Pr \left( \xi_{t',n}^\text{pres}(I) \forall t' < t, \xi_{t,n}^\text{pres} \leq \tilde{c} \right) \leq w_{t}^\text{pres}(\bar{I}) \alpha. \quad (5)$$

The significance threshold $c_{a_t}(I)$ for testing $H_{a_t}$ is defined as the largest $\tilde{c}$ such that

$$\Pr \left( \xi_{t',n}^\text{pres}(I) \forall t' \leq t, p_{a_t} \leq \tilde{c} | H_{a_t} \right) \leq w_{a_t}(\bar{I}) \alpha. \quad (6)$$

Crucially, these calculations do not depend on observing counterfactual or future hypotheses. Using a union bound, we can show that the Type I error for falsely rejecting the intersection hypothesis $I$ using the critical values defined above is bounded by the sum of the right hand sides of (5) and (6) for all $(t, a_t)$ in $I$. Because the total weight in the graph is always one, we achieve Type I error control at level $\alpha$. Using this idea, we can show that this SRGP indeed controls the FWER:

**Theorem 2.** Algorithm using significance thresholds defined using equations (5) and (6) control FWER at level $\alpha$ for adaptively selected hypotheses.

3 SIMULATION STUDIES

We now present two simulation studies of model developers who adaptively propose modifications to their initial ML algorithm. The developers aim to improve the model’s area under the receiver operating characteristic curve (AUC) and quantify the performance increase as accurately as possible. Because our adaptive test data reuse procedures only release a single bit of information at each iteration, we must carefully design the hypothesis tests to obtain a numeric bound on the performance improvement. In particular, we define the $j$-th adaptive hypothesis test as

$$H_{0,j}^\text{adapt}: \psi\left(\hat{f}_j^\text{adapt}, P_0\right) \leq \psi\left(\hat{f}_0; P_0\right) + \delta_j^\text{adapt} \quad (7)$$

where $\psi(f, P)$ denotes AUC of model $f$ for distribution $P$ and $\delta_j^\text{adapt} \geq 0$ is the improvement difference that we are trying to detect. To ensure the model performance tends to improve with each approval, we set $\delta_j^\text{adapt} = \delta_j^\text{adapt} + 0.01$ whenever the $j$-th null hypothesis is rejected. Note that one could consider more complicated hypotheses, each with their pros and cons. For example, one can check that the modifications are strictly improving and test for an improvement difference; however, this can be overly stringent.

The purpose of the first simulation study is to investigate FWER control. We do this by simulating a model developer who tries to overfit to the test data based on the information released at each iteration. The purpose of the second simulation study is to investigate power. Here the model developer generally proposes good algorithmic modifications by continually refitting the model given an IID data stream.

In both simulations, we generate $X \in \mathbb{R}^{100}$ using a multivariate Gaussian distribution. $Y$ is generated using a logistic regression model where the coefficients of the first six variables are 0.75 and all other model parameter are zero. The modifications are also logistic regression models. We evaluate the two SRGPs proposed in this paper—SRGP with fixed sequence tests (fsSRGP) and SRGP with hypothetical prespecified model updates (presSRGP)—against relevant baseline comparators, including the standard Bonferroni procedure (Bonferroni) and the Bonferroni-based SRGP (bonfSRGP). The weights in the SRGPs were defined such that the first outgoing edge (a successful rejection of the hypothesis) is 0.8 and for each subsequent edge, it was assigned 0.8 of the remaining weight. Unless specified otherwise, all the MTPs control the FWER at level $\alpha = 0.1$. Details for deriving test statistics and significance thresholds are provided in the Appendix.

3.1 VERIFYING FWER CONTROL

Here we show how MTPs that fail to control the FWER can drastically elevate one’s risk of overfitting to the test data, as compared to appropriately-designed adaptive test data reuse procedures. In particular, we consider the naïve procedure that tests every adaptive hypothesis at level $\alpha$ (BinaryThres). The reusable test dataset has 100 obser-
vations and the model developer tests $T = 50$ modifications. For the purpose of illustration, the initial model is set to the oracle, so all proposed modifications are unacceptable.

The simulated model developer tries to find models that overfit to the test data by searching within the neighborhood of the currently approved model. In particular, the developer iteratively perturbs the coefficient of each irrelevant variable by 0.6 in the positive and negative directions. When any such modification is approved, the model developer will continue perturbing that coefficient in the same direction until it fails to reject the null hypothesis. For presSRGP, the prespecified model update at iteration $t$ is the model with coefficients exactly the same as the initial model except that the coefficient for the $(7 + \lfloor t/2 \rfloor)$-th variable is set to 0.6 if $t$ is even and -0.6 if $t$ is odd.

Figure 3 shows the result from 400 replicates. Notably, BinaryThres approves at least one inferior modification with probability 75% and concludes that the modifications by the last iteration improves the AUC by at least 0.02, even though the AUC actually drops by 0.025 on average. All the other MTPs appropriately control the FWER at the desired rate of 10% and, thus, protect against over-fitting.

### 3.2 ASSESSING POWER

Here the simulated model developer has access to an IID data stream and iteratively refits a logistic regression model on this data. Because training on more data from the target population tends to improve model performance, the modifications are usually beneficial. However, there is a risk that the modification does not improve performance or that the improvement is negligible, especially because there is a potential for overfitting to the reusable test data set. By testing hypotheses \( H_0 \), we can restrict approval to only those model updates with meaningful improvements in the AUC.

The test dataset has 800 observations and we allow $T = 15$ adaptive tests. At each time point, the model developer receives a new observation and refits the model. To spend alpha more judiciously, the model developer will only submit the refitted model if the power calculations suggest that the probability for rejecting the null hypothesis exceeds 50%. Specifically, they perform power calculations by setting the true performance improvement to the CI lower bound, which is estimated using split-sample validation. (For simplicity, the power calculations do not perform any multiple testing correction.) To run presSRGP, the prespecified model updating procedure also selects updates based on a hypothesis test similar to \( H_0 \) but replacing the adaptive difference sequence $\delta_j^{\text{adapt}}$ with the prespecified difference sequence $\delta_j^{\text{pres}} = 0.0025(j - 1)$ as well as replacing the adaptive modifications with the prespecified ones.

The procedures differed significantly in power (Figure 4). On average, Bonferroni approved two modification, BonfSRGP approved 4.5, fsSRGP approved 5, and presSRGP approved 5.5. By the end of the testing procedure, the average AUC of the approved model by presSRGP was 0.8 whereas Bonferroni only attained an AUC of 0.75. Finally, the detected performance improvement was highest using presSRGP, as compared to the other methods.

### 4 REVISING PREDICTIONS FOR ACUTE HYPTENSION EPISODES

We now apply our procedure for approving modifications to a risk prediction model for acute hypotension episodes (AHEs), one of the most frequent critical events in the intensive care unit (ICU) (Walsh et al. 2013). The ICU is a clinical environment that continuously generates high throughput data. Thus, a model developer can readily collect new data in this setting to retrain an existing model. To mimic this, we use data from the eICU Collaborative Research Database (Pollard et al. 2018). We train an initial model on 40 randomly selected admissions and simulate a data stream in which a randomly selected admission is observed at each time point. The reusable test data is composed of 500 admissions and is used to evaluate $T = 15$ modifications.

The task is to predict AHE 30 minutes in advance, where we define AHE as any 5-minute time period where the average mean arterial pressure (MAP) falls below 65 mmHg. The input features to the model are baseline variables age, sex, height, and weight; vital signs MAP, heart rate, and respiration rate at the current time point; and the same set of vital signs five minutes prior. The prediction model is a gradient boosted tree (GBT) and is continually refit on the incoming data.

Here we consider a more complex hypothesis test that checks for calibration-in-the-large (Steyerberg 2009) and improvement in AUC. The $j$-th adaptive null hypothesis is

\[
H_{0,j}^{\text{adapt}} : \psi \left( \hat{f}^{\text{adapt}}_j ; P_0 \right) - \psi \left( \hat{f}_0 ; P_0 \right) + \delta_j^{\text{adapt}} \leq \epsilon, \quad \text{or} \quad E \left[ \hat{f}^{\text{adapt}}_j (X) - Y \right] \notin [-\epsilon, \epsilon],
\]

where $\delta_j^{\text{adapt}}$ is defined using the same procedure as that in Section 3.2 $\hat{f}_j$ is the modification determined to have
sufficient power for rejecting the null, and margin of error \( \epsilon \) is 0.05. We will refer to \( E \left[ \hat{f}_{\text{adapt}}(X) - Y \right] \) as calibration-error-in-the-large. Details on calculating the test statistic and significance thresholds are provided in the Appendix.

Results from 40 replicates are shown in Figure 5. We observe the same ranking of MTPs as that in Section 3.2: prespecSRGP performed the best, followed by fsSRGP. Compared to the previous section, the relative improvement between the methods is smaller because the GBTs improved rapidly at early time points and slowed down thereafter.

5 RELATED WORK

Our paper relates to a large body of work on methods for providing valid statistical inference and preventing false discoveries. Much of this literature has focused on testing prespecified hypotheses on the same dataset while controlling the FWER (Hochberg and Tamhane, 1987; Westfall et al., 2010), false discovery rate (FDR) (Benjamini and Hochberg, 1995), or some variant thereof (van der Laan et al., 2004). More recent works consider testing a sequence of adaptive hypotheses on prospectively-collected data from a data stream and controlling online error rates (Foster and Stine, 2008; Ramdas et al., 2018). This work considers the setting where we adaptively test hypotheses on the same dataset. To control the bias, testing procedures must limit the amount of information released about the test dataset (Russo and Zou, 2016). Techniques based on differential privacy, which is a mathematically rigorous formalization of data privacy (Dwork and Roth, 2014), do this by adding random noise (e.g., Laplace or Gaussian noise) to the test statistic or, more generally, the queried result (Dwork et al., 2015a). While theoretical guarantees are available for differential privacy based methods for test data reuse (e.g., (Dwork et al., 2015b; Russo and Zou, 2016), Rogers et al., 2016; Cummings et al., 2016; Dwork et al., 2017; Feldman and Steinke, 2017, 2018; Shenfeld and Ligett, 2019; Gossmann et al., 2021 and others), the required size of the test dataset is prohibitively large for many application domains or require injecting very large amounts of noise (Rogers et al., 2019; Gossmann et al., 2021). An alternative approach is to directly limit the number of bits of information released to the model developer by discretizing the queried result along some grid (Blum and Hardt, 2015). Existing methods essentially perform a Bonferroni correction for the number of distinct hypotheses, which also require unreasonably large test datasets for many applications. To improve testing power, a number of works have assumed that the adaptivity of the model developer is limited (e.g., the models are highly correlated, or the model developer is not entirely “adversarial”) to justify the use of a less conservative correction factor (Mania et al., 2019; Zrnic and Hardt, 2019). In contrast, the SRGPs proposed in this work achieve higher power via alpha-recycling and account for the correlation structure without needing to make assumptions about the model developer.

6 CONCLUSION

We show how to leverage SRGPs to design valid and powerful approaches for testing a sequence of adaptively-defined algorithmic modifications on the same dataset. The overall steps of this framework are (i) limit the amount of information leakage by reporting only binary test results (approve versus deny modifications), (ii) spend and recycle alpha using an SRGP, and (iii) design consonant, closed-testing procedures whose significance thresholds can be computed without needing to observe the counterfactual hypotheses. To account for correlation between the algorithmic modifications, we presented two new SRGPs. fsSRGP achieves higher power by leveraging the correlation structure between the observed algorithmic modifications. presSRGP asks the model developer to generate a sequence of algorithmic modifications using a prespecified learning procedure and leverages the correlation between the adaptive and prespecified algorithmic modifications. In empirical studies, these procedures approved more algorithmic modifications than existing methods, with presSRGP achieving the highest power.

One direction of future work is to optimize the power for approving algorithmic modifications by (i) tuning the node and edge weights in the SRGP and (ii) exploring various testing strategies that the model developer can employ. In addition, model developers are often interested in obtaining more detailed test results like p-values and confidence intervals. Another direction for future work is to design SRGPs that release more information per iteration, perhaps by leveraging differential privacy techniques.
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A PROOFS

Lemma 1. The adaptive SRGP in Algorithm 1 with a fixed strategy is equivalent to a prespecified SRGP.

Proof. We define a filtration over approval histories up to the maximum number of iterations $T$. That is, define sample space $\Omega$ as the set of approval histories over $T$ iterations, i.e. $\Omega = \{0, 1\}^{T-1}$, and $\sigma$-algebras $\mathcal{F}_t$ for $t = 1, \ldots, T$ over approval histories up to iteration $t$. To show that the adaptive SRGP is equivalent to a prespecified SRGP, we need to show that the adaptive procedure defines a set of hypotheses, node weights, and edge weights for the initial set of hypotheses $I_0$, the hypotheses and weights are $\mathcal{F}_1$-measurable functions, and the weight constraints are satisfied. First, we note that the edge weights being elicited at iteration $t$ in Algorithm 1 is equivalent to eliciting the edge weights for the initial set of hypotheses $I_0$, i.e. $g_{a_{t-1}, a_t} = g_{a_{t-1}, a_t}(I_0)$ in Algorithm 1. This is because we only elicit the edge weight $g_{a_{t-1}, a_t}$ if there has been no approval since time $t_2$ so the edge weights being elicited are never updated via the edge-weight renormalization step in SRGs. As such, the adaptive SRGP in Algorithm 1 for a model developer with a fixed strategy for selecting hypotheses and weights can be described to have a fixed hypothesis testing tree structure with

- $\mathcal{F}_t$-measurable hypotheses $H_{a_t}(I_0)$ for all $a_t$
- $\mathcal{F}_1$-measurable node weights $w_{a_t}(I_0)$ for all $a_t \in \{0, 1\}^{T-1}$ that satisfy the constraint that they sum to one,
- and $\mathcal{F}_t$-measurable edge weights $g_{a_{t-1}, a_t}(I_0)$ for all valid edges $(a_{t-1}, a_t)$ in the graph that satisfy the constraint that all outgoing edge weights sum to one.

Although the hypotheses and edge weights are $\mathcal{F}_t$-measurable, they can also be viewed as $\mathcal{F}_1$-measurable functions over the input space $a_t$ and $(a_{t-1}, a_t)$, respectively. Moreover, the edge weights satisfy the edge weight constraints by design. Thus the adaptive SRGP satisfies the node and edge weights constraints with respect to $\mathcal{F}_1$.

Lemma 2. If the adaptive SRGP in Algorithm 1 controls the FWER for any fixed strategy, then the adaptive SRGP in Algorithm 1 controls the FWER for any stochastic strategy.

Proof. Let $S$ be the set of all fixed strategies. The stochastic adaptive strategy is a random distribution over $S$. Its FWER is

$$\Pr\left(\text{incorrectly reject some } H_t^{\text{adapt}}\right) = \sum_{s \in S} \Pr(S = s) \Pr\left(\text{incorrectly reject some } H_t^{\text{adapt}} \mid S = s\right)$$

where the latter probability on the right hand side is the FWER for a fixed strategy $s$. As such, the FWER of the stochastic strategy is properly controlled as long as the FWER of any fixed strategy is properly controlled.

Corollary 1. Algorithm 1 with the significance thresholds defined per

$$c_{a_t}(I_t) = w_{a_t}(I_t)\alpha$$

controls the FWER at level $\alpha$.

Proof. Per Lemmas 1 and 2, it suffices to show that the fully prespecified SRGP controls the FWER. Recall that (9) is a closed weighted Bonferroni test in Bretz et al. (2011a). As such, any fixed or stochastic adaptive strategy would control FWER.

Proof for Theorem 1. Per Lemmas 1 and 2, it suffices to show that the fully prespecified SRGP controls the FWER.

First, per the proof in Bretz et al. (2009), we note that node weights for any intersection hypothesis $I$ calculated using Algorithm 1 are well-defined, in that it does not depend on ordering in which we remove nodes from the graph.

We begin with proving that for any intersection hypothesis $I$, the critical values calculated using (4) controls the Type I error. First we show that for any $a_t$ ending with success (i.e. $a_{t,t-1} = 1$) and any $I$, the calculated critical values for testing the
We first prove that the critical values per (6) control the Type I error for any intersection hypothesis \( \cap_{a_k \in G_{a_i} \cap I} H_{a_k} \) controls the Type I error at level \( \left( \sum_{a_k \in G_{a_i} \cap I} w_{a_k}(I) \right) \alpha \). Per the definition of the critical values in (4), we have that

\[
\Pr \left( \text{we reject for some } a_j \in G_{a_i} \cap I \mid H_{G_{a_i} \cap I} \right) = \sum_{a_j \in (G_{a_i} \cap I)} \Pr \left( p_{a_k} > c_{a_k}(I) \forall a_k \in G_{a_i} \cap I, k < j, p_{a_j} < c_{a_j}(I) \mid \cap_{a_k \in G_{a_i} \cap I, k \leq j} H_{a_k} \right) \leq \left( \sum_{a_j \in (G_{a_i} \cap I)} w_{a_j}(I) \right) \alpha.
\]

Therefore, as long as the total node weight across \( I \) is no more than one, we control the Type I error at level \( \alpha \). Because Type I error control holds for all intersection hypotheses \( I \), we have established that this procedure is a valid closed test.

Next, per the proof in Bretz et al. (2009), we must show that the critical values satisfy the monotonicity condition to prove that our procedure is a valid consonant, shortcut procedure. More specifically, we require the following to hold for all \( t = 1, \ldots, T \):

\[
c_{a_t}(I) < c_{a_t}(J) \quad \forall J \subseteq I.
\]

The proof is by induction. It is easy to see that (10) holds for \( t = 1 \). Suppose (10) holds for \( 1, \ldots, t - 1 \). Now consider any history \( a_t \) that ends with an approval. Consider any \( a_t \) and subset \( J \subseteq I \) such that \( a_t \in G_{a_i} \cap J \). We have that

\[
c_{a_t}(J) = \sup \left\{ \hat{c} : \Pr \left( p_{a_k} > c_{a_k}(J) \forall a_k \in K, p_t \in \hat{c} \mid H_{K \cup \{a_t\}} \right) \leq \left( \sum_{a_k \in (G_{a_i} \cap J) \setminus K} w_{a_k}(J) \right) \alpha \forall K \subseteq \{ a_k : a_k \in G_{a_i} \cap J, k < t \} \right\}
\]

\[
\geq \sup \left\{ \hat{c} : \Pr \left( p_{a_k} > c_{a_k}(I) \forall a_k \in K, p_t \in \hat{c} \mid H_{K \cup \{a_t\}} \right) \leq \left( \sum_{a_k \in (G_{a_i} \cap J) \setminus K} w_{a_k}(I) \right) \alpha \forall K \subseteq \{ a_k : a_k \in G_{a_i} \cap J, k < t \} \right\}
\]

\[
= c_{a_t}(I)
\]

where the first inequality follows by induction and the second inequality is because the weights are monotonic.

**Proof for Theorem 2** Per Lemmas 1 and 3 it suffices to show that the fully prespecified SRGP controls the FWER.

We first prove that the critical values per (6) control the Type I error for any intersection hypothesis \( I \). For any \( I \), define \( \hat{I} \) as the union of \( I \) and all prespecified nodes. Then the Type I error can be bounded using a sequence of union bounds:

\[
\Pr \left( \exists (t, a_t) \in I \text{ s.t. } p_{a_t} < c_{a_t}(I) \mid H_I \right) \leq \Pr \left( \exists t \text{ s.t. } \xi_{t,n}^{\text{pres}} \leq \xi_{t,n}^{\text{pres}}(I) \right) \text{ OR } \exists (t, a_t) \in I \text{ s.t. } p_{a_t} < c_{a_t}(I) \mid H_I \right)
\]

\[
\leq \sum_{t=1}^{\infty} \Pr \left( \xi_{t,n}^{\text{pres}} > \xi_{t,n}^{\text{pres}}(I) \forall t' \leq t - 1, \xi_{t,n}^{\text{pres}} \leq \xi_{t,n}^{\text{pres}}(I) \mid H_I \right) + \sum_{a_t \in \hat{I}} \Pr \left( \xi_{t,n}^{\text{pres}} > \xi_{t,n}^{\text{pres}}(I) \forall t' \leq t, p_{a_t} < c_{a_t}(I) \mid H_I \right)
\]

\[
\leq \left( \sum_{t=1}^{\infty} \left( w_{t}^{\text{pres}}(\hat{I}) + \sum_{a_t \in \hat{I}} w_{a_t}(\hat{I}) \right) \right) \alpha
\]

\[= \alpha.
\]

Because the weights are nondecreasing in Algorithm 1, the critical values defined in (6) satisfy the monotonicity condition. As such, Algorithm 1 is a consonant, short-cut procedure for the above closed test.
B HYPOTHESIS TEST DETAILS

B.1 TESTING FOR AN IMPROVEMENT IN AUC

In Section 3, we decide whether or not to approve a modification by testing the adaptively-defined null hypothesis $H_{0,a_j}$ at each iteration $j$, which compares the AUC between the $j$th adaptively proposed model and the initial model. Per Algorithm 1, we test the adaptive hypotheses by treating them as pre-specified hypotheses from a bifurcating tree, i.e.

$$H_{0,a_j} : \psi(\hat{f}_{a_j}, P_0) \leq \psi(\tilde{f}_0; P_0) + \delta_{a_j} \tag{11}$$

for approval histories $a_j$. We now describe how the test statistics and significance thresholds are constructed.

Recall that the AUC is equal to the Mann-Whitney U-statistic for comparing ranks across two populations, i.e.

$$\psi(f, P_0) = P_0 \left( f(X_1) > f(X_2) \mid Y_1 = 1, Y_2 = 0 \right), \tag{12}$$

where $(X_1, Y_1)$ and $(X_2, Y_2)$ represent independent draws from $P_0$. The empirical AUC is defined as

$$\psi(f, P_n) = \frac{1}{n_0n_1} \sum_{i=1}^{n_0} \sum_{j=1}^{n_1} \mathbb{1}\{f(X_j) > f(X_i)\} \mathbb{1}\{Y_j = 1, Y_i = 0\}, \tag{13}$$

where $n_0$ is the number of observations with $Y = 0$ and $n_1 = n - n_0$. To test (11), we characterize the asymptotic distribution of (13) by analyzing its influence function. Given IID observations from $P_0$, (13) is an asymptotically linear estimator of the model’s AUC (LeDell et al., 2015), in that

$$\psi(f, P_n) - \psi(f, P_0) = \frac{1}{n} \sum_{i=1}^{n} \phi(f, P_0)(X_i, Y_i) + o_p(1/\sqrt{n}) \tag{14}$$

with influence function

$$\phi(f, P_0)(X_i, Y_i) = \frac{\mathbb{1}\{Y_i = 1\}}{P_0(Y = 1)} P_0(f(X) < c \mid Y = 0; c = f(X_i))$$

$$+ \frac{\mathbb{1}\{Y_i = 0\}}{P_0(Y = 0)} P_0(f(X) > c \mid Y = 1; c = f(X_i))$$

$$- \left\{ \frac{\mathbb{1}\{Y_i = 0\}}{P_0(Y = 0)} + \frac{\mathbb{1}\{Y_i = 0\}}{P_0(Y = 0)} \right\} \psi(f, P_0).$$

Per the Central Limit Theorem, we have that

$$\sqrt{n} \left( \psi(f, P_n) - \psi(f, P_0) \right) \rightarrow_d N \left( 0, \sigma(f, P_0)^2 \right) \tag{15}$$

where $\sigma(f, P_0)^2 = \text{Var}(\phi(f, P_0)(X, Y))$. We can then test the null hypothesis $H_0 : \psi(\tilde{f}_0, P_0) \leq c$ for some constant $c$ based on the asymptotic normality of (13). In addition, we can test (11) by deriving the asymptotic distribution of $\psi\left(\hat{f}_{a_j}, P_0\right) - \psi\left(\tilde{f}_0; P_0\right)$ based on the difference of the influence functions $\phi(\hat{f}_{a_j}, P_0)(X, Y) - \phi(\tilde{f}_0, P_0)(X, Y)$. To run fsSRGP, we can extend the above derivations to construct a fixed sequence test for testing a family of null hypotheses (11) across multiple iterations $j$ by analyzing the joint asymptotic distribution of the test statistics $\psi\left(\hat{f}_{a_j}, P_n\right) - \psi\left(\tilde{f}_0; P_n\right)$ and compute the significance thresholds defined in (4). Similar logic can be used to derive the critical values (5) and significance thresholds (6) in fsSRGP.

B.2 TESTING MODEL DISCRIMINATION AND CALIBRATION

Section 4 considers the more complex hypothesis test (8), which checks for an improvement in AUC and calibration-in-the-large. We implement this by testing three individual hypothesis tests using sequential gatekeeping. First, we test that the difference between the average risk prediction and the observed event rate is no smaller than $-\epsilon$. Next, we test that this difference is no larger than $\epsilon$. Finally, we test for an improvement in AUC using the procedure described in Section B.1. To control the Type I error for rejecting the overall null hypothesis, we perform alpha spending across the individual hypotheses.