# Better-Than-Chance Classification for Signal Detection

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1 Abstract

[TODO]

#### 1 Introduction

4 A common workflow in neuroimaging consists of fitting a classifier, and es-

5 timating its predictive accuracy using cross validation. Given that the cross

validated accuracy is a random quantity, it is then common to test if the

cross validated accuracy is significantly better than chance using a permu-

tation test. Examples in the neuroscientific literature include Golland and

Fischl [2003], Pereira et al. [2009], Varoquaux et al. [2016], and especially

the recently popularized multivariate pattern analysis (MVPA) framework of Kriegeskorte et al. [2006]. This practice is also observed in very high profile publications in the genetics literature: Golub et al. [1999], Slonim et al.

[2000], Radmacher et al. [2002], Mukherjee et al. [2003], Juan and Iba [2004],

Jiang et al. [2008].

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To fix ideas, we will adhere to a concrete example. In Gilron et al. [2016], the authors seek to detect brain regions which encode differences between vocal and non-vocal stimuli. Following the MVPA workflow, the localization problem is cast as a supervised learning problem: if the type of the stimulus can be predicted from the spatial activation pattern significantly better than chance, then a region is declared to encode vocal/non-vocal information. We call this an accuracy test, a.k.a. class prediction, or pattern discrimination

This same signal detection task can be also approached as a two-group multivariate test. Inferring that a region encodes vocal/non-vocal information, is essentially inferring that the spatial distribution of brain activations is different given a vocal/non-vocal stimulus. As put in Pereira et al. [2009]:

... the problem of deciding whether the classifier learned to discriminate the classes can be subsumed into the more general question as to whether there is evidence that the underlying distributions of each class are equal or not.

A practitioner may then call upon a two-group population test such as Hotelling's  $T^2$  [Anderson, 2003]. Alternatively, if the size of a brain region is large compared to the number of observations, so that the spatial covariance cannot be fully estimated, then a high dimensional version of Hotelling's test can be called upon, such as in Schäfer and Strimmer [2005] or Srivastava [2007]. For brevity, and in contrast to accuracy tests, we will call any two-sample multivariate tests simply population tests, also termed class comparisons. [TODO: rename to parameter test?]

At this point, it becomes unclear which is preferable: a population test or an accuracy test? The former with a heritage dating back to Hotelling [1931], and the latter being extremely popular, as the 959 citations<sup>1</sup> of Kriegeskorte et al. [2006] suggest.

The comparison between location and accuracy tests was precisely the goal of Ramdas et al. [2016], who compared the  $T^2$  population test to the accuracy of Fisher's linear discriminant analysis classifier (LDA). By comparing the rates of convergence of the powers to 1, Ramdas et al. [2016] concluded that accuracy and population tests are rate equivalent.

Asymptotic relative efficiency measures (ARE) are typically used by statisticians to compare between rate-equivalent test statistics [van der Vaart, 1998]. Ramdas et al. [2016] derive the asymptotic power functions of the two test statistics, which allows to compute the ARE between Hotelling's  $T^2$  (location) test and Fisher's LDA (accuracy) test. Theorem 14.7 of van der Vaart [1998] relates asymptotic power functions to ARE. Using the results of Ramdas et al. [2016] we deduce that the ARE is lower bounded by  $2\pi \approx 6.3$ . This means that Fisher's LDA requires at least 6.3 more samples to achieve the same (asymptotic) power than the  $T^2$  test. In this light, the accuracy test is remarkably inefficient compared to the population test. For comparison, the t-test is only 1.04 more (asymptotically) efficient than Wilcoxon's rank-sum test [Lehmann, 2009], so that an ARE of 6.3 is strong evidence in favor of the population test.

Before discarding accuracy tests as inefficient, we recall that Ramdas et al. [2016] analyzed a *half-sample* holdout. The authors conjectured that a leave-one-out approach, which makes more efficient use of the data, may have better performance. Also, the analysis in Ramdas et al. [2016] is asymptotic. This eschews the discrete nature of the accuracy statistic, which will be

<sup>&</sup>lt;sup>1</sup>GoogleScholar. Accessed on Aug 4, 2016.

shown to have crucial impact. Since typical sample sizes in neuroscience are not large, we seek to study which test is to be preferred in finite samples? Our conclusion will be quite simple: population tests almost always have more power than accuracy tests.

Our statement rests upon the observation that with typical sample sizes, the accuracy test statistic is highly discrete. Permutation testing with discrete test statistics are known to be conservative [Hemerik and Goeman, 2014], since they are insensitive to mild perturbations of the data, and they cannot exhaust the permissible false positive rate. The degree of discretization is governed by the number of samples. In our neuroscience example from Gilron et al. [2016], the classification is performed based on 40 trials, so that the test statistic may assume only 40 possible values. This number of examples is not unusual if considering this is the number of trial-repeats, or the number of subjects in an neuroimaging study.

The discretization effect is aggravated if the test statistic is highly concentrated. For an intuition consider the usage of a the resubstitution accuracy as a test statistic. This statistic simply means that the accuracy is not cross validated. If the data is high dimensional, the resubstitution accuracy will be very high due to over fitting. In a very high dimensional model, the resubstitution accuracy will be 1 for the observed data [McLachlan, 1976, Theorem 1], but also for any permutation. The concentration of resubstitution accuracy near 1, and its discreteness, render this test completely useless, with a power tending to 0 for any (fixed) effect size, as the dimension of the model grows.

To compare the power of accuracy tests and population tests in finite samples, we perform a simulation study of a battery of test statistics. We start with formalizing the problem in Section 2. The main findings are reported in Sections 4 and 5. A discussion follows in Section 6.

## <sup>93</sup> 2 Problem setup

Let  $y \in \mathcal{Y}$  be a class encoding. Let  $x \in \mathcal{X}$  be a p dimensional feature vector. In our vocal/non-vocal example we have  $\mathcal{Y} = \{-1, 1\}$  and p, the number of voxels in a brain region so that  $\mathcal{X} = \mathbb{R}^{27}$ .

Given n pairs of  $(x_i, y_i)$ , typically assumed i.i.d., a population test amounts to testing whether x|y=1 has the same distribution as x|y=-1. I.e., we test if the multivariate voxel activation pattern has the same distribution when given a vocal stimulus, as when given a non-vocal stimulus.

An accuracy test amounts to learning a predictive model and testing if its predictions are better than chance. Denoting a dataset by  $\mathcal{S} := (x_i, y_i)_{i=1}^n$ ,

and a predictor by f, then a learning algorithm  $\mathcal{A}$  is a mapping  $\mathcal{A}: \mathcal{S} \to f$ . The accuracy of predictor  $f = \mathcal{A}_{\mathcal{S}}$  is defined as the probability of f making a correct prediction. Denoting by P the probability measure of  $(\mathbf{x}, \mathbf{y})$ , and by  $P^n$  the same for the i.i.d sample  $\mathcal{S}$ 

$$\mathcal{E}_{\mathcal{A}_{\mathcal{S}}} := P(\mathcal{A}_{\mathcal{S}}(x) = y). \tag{1}$$

The accuracy of an algorithm  $\mathcal{A}$  is defined as the average accuracy, over all possible data sets

$$\mathcal{E}_{\mathcal{A}} := \int_{\mathcal{S}} \mathcal{E}_{\mathcal{A}_{\mathcal{S}}} dP^{n}(\mathcal{S}). \tag{2}$$

Denoting an estimate of  $\mathcal{E}_f$  by  $\hat{\mathcal{E}}_f$ , a statistically significant "better than chance" estimate of  $\hat{\mathcal{E}}_f$  is evidence that the classes are distinct.

#### 2.1 Candidate Tests

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The design of a permutation test using  $\hat{\mathcal{E}}_f$ , requires the following design choices:

- 1. Is  $\hat{\mathcal{E}}_f$  cross validated or not?
- 2. For a V-fold cross validated test statistic:
  - (a) Should the data be refolded in each permutation?
  - (b) Should the data folding be balanced (a.k.a. stratified)?
  - (c) How many folds?
- 3. How to estimate  $\hat{\mathcal{E}}_f$ ?

We will now address these questions while bearing in mind that unlike the typical supervised learning setup, we are not interested in an unbiased estimate of  $\hat{\mathcal{E}}_f$ , but rather in the mere detection of a difference between two classes.

Cross validate or not? Given our goal, a biased error estimate is not a problem provided that bias is consistent over all permutations. The underlying intuition is that if the exact same computation is performed over all permutations, then a permutation test will be "fair", i.e., will not inflate the false positive rate. We will thus be considering both cross validated accuracies, and *resubstitution accuracies*, where the accuracy is evaluated on the training set and not on a holdout.

Balanced folding? The standard practice when cross validating is to constrain the data folds to be balanced, i.e. stratified [e.g. Ojala and Garriga, 2010]. This means that each fold has the same number of examples from each class. We will report results with both balanced and unbalanced data foldings, only to discover, it does not really matter.

Refolding? The standard practice in neuroimaging is to permute labels and refold the data after each permutation, so that the balance of the classes in each fold is preserved. We will adhere to this practice due to its popularity, even though it can be simplified by permuting features instead of labels, as done by Golland et al. [2005].

How many folds? Different authors suggest different rules for the number of folds. We will be varying the number of folds, and ultimately discover that the power decreases with the number of folds.

How to estimate accuracy? Given a predictor for the i'th data example<sup>2</sup>,  $f_i$ , a natural test statistic its empirical accuracy

$$\hat{\mathcal{E}}_f := \sum_i I(f_i(x_i) = y_i). \tag{3}$$

Since low accuracies, even 0, are evidence that the classes are separated, we can consider the departure from chance level  $|\hat{\mathcal{E}}_f - 0.5|$ , as another candidate test statistic. For unbalanced classes, chance level is not 0.5, but rather the the probability of the majority class, we denote by  $\hat{\pi}$ . This suggests the following test statistic  $|\hat{\mathcal{E}}_f - \hat{\pi}|$ . Since we will be aggregating these statistics over random data sets where  $\hat{\pi}$  may vary, it seems appropriate to standardize the scale. We thus propose the z-scored accuracy statistic,  $\hat{\mathcal{Z}}_f$ :

$$\hat{\mathcal{Z}}_f := |\hat{\mathcal{E}}_f - \hat{\pi}| / \sqrt{\hat{\pi}(1 - \hat{\pi})}. \tag{4}$$

Table 1 collects an initial battery of tests we will be comparing.

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<sup>&</sup>lt;sup>2</sup>We add the unorthodox observation index i to  $f_i$  to accommodate for cross validation methods, where each prediction is made with a different classifier, fit to a slightly different dataset.

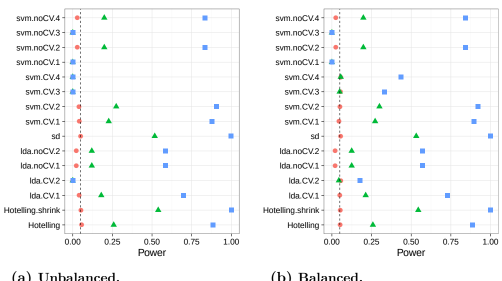
Name	Basis	CV	Accuracy	Parameters
Hotelling	Hotelling	_	_	_
Hotelling.shrink	Hotelling	_	_	_
lda.CV.1	LDA	V-fold	$\hat{\mathcal{E}}$	_
lda.CV.2	LDA	V-fold	$\hat{\mathcal{Z}}$	_
lda.noCV.1	LDA	_	accuracy	_
lda.noCV.2	LDA	_	z-accuracy	_
$\operatorname{sd}$	SD	_	_	_
svm.CV.1	SVM	V-fold	accuracy	cost=1e1
svm.CV.2	SVM	V-fold	accuracy	cost=1e-1
svm.CV.3	SVM	V-fold	z-accuracy	cost=1e1
svm.CV.4	SVM	V-fold	z-accuracy	cost=1e-1
svm.noCV.1	SVM	_	accuracy	cost=1e1
svm.noCV.2	SVM	_	accuracy	cost=1e-1
svm.noCV.3	SVM	_	z-accuracy	cost=1e1
svm.noCV.4	SVM	_	z-accuracy	cost=1e-1

Table 1: This table collects the various test statistics we will be studying. Three are population tests: Hotelling, Hotelling.shrink, and sd. Hotelling is the classical two-group  $T^2$  statistic. Hotelling.shrink is a high dimensional version with the regularized covariance in Schäfer and Strimmer [2005]. sd is another high dimensional version of the  $T^2$ , from Srivastava et al. [2013]. The rest of the tests are variations of the linear SVM, and Fisher's LDA, with varying accuracy measures, cross validated or not, and varying tuning parameters. For example, svm.CV.4 is a linear SVM implemented with the svm R function, the cost parameter set at 0.1, and using the cross validated z-scored accuracy in Eq. 4. Another example is lda.noCV.1, which is Fisher's LDA, returning the resubstitution accuracy.

## 3 Controlling the False Positive Rate

Figure 1 demonstrates that all of the tests considered conserve the desired 0.05 false positive rate, up to varying levels of conservatism. This can be seen from the fact that the probability of rejection is no larger than 0.05 in the absence of any effect, encoded by a red circle. This is true, in particular if: (a) the folds are balanced or not, (b) the tuning parameters of some test statistic are varied, (d) the number of folds is varied. We also observe that the most conservative tests are the resubstitution accuracy statistics. We return to this matter in the Discussion.

Figure 1: The power of a permutation test with various test statistics. The power on the x axis. Effect are color and shape coded. The various statistics on the y axis. Their details are given in Table 1. Effects vary over 0 (red circle), 0.25 (green triangle), and 0.5 (blue square). Simulation details in Appendix B. Cross-validation was performed with balanced and unbalanced data folding. See sub-captions.



(a) Unbalanced.

(b) Balanced.

#### 4 Power

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Having established that all of the tests in our battery control the false positive rate, it remains to be seen if they have similar power- especially when comparing population tests to accuracy tests. From the simulation results reported in Appendix C we collect the following insights:

- 1. population tests have more power than accuracy tests in all our configurations.
- 2. The conservativeness decays as the sample grows (Figures 9a, 9b and
  - 3. For heavy tailed distributions (Figure 8b), the extra power of the location test vanishes.
  - 4. The presence of correlations between coordinates reduces the signal to noise ratio (SNR), thus reduces power. More importantly, in the presence of correlations the effect of regularization is amplified, increasing the power difference between regularized and non-regularized test

- statistics. Put differently- in low SNR regimes, regularization proves crucial (Figure 10b).
- 5. The z-scoring of the accuracies was introduced to deal with unbalanced foldings. If the z-scoring has any effect at all, it merely kills power.
- 6. Both accuracy and population tests are inappropriate for scale alternatives (Figure 8a). This was to be expected and is reported mostly as a sanity check.
  - 7. Balanced folding only affects the z-scored accuracy, in the opposite direction than we anticipated.
  - 8. Increasing the SVM's cost parameter, which reduces the number of support vectors entering the classifier, reduces power.

The major insight from simulations is that the use of accuracy tests for signal detection is underpowered compared to population tests. We now verify this finding on a neuroimaging dataset.

## 5 Neuroimaging Example

Figure 2 is an application of both a location and an accuracy test to the data of Pernet et al. [2015]. The authors of Pernet et al. [2015] collected fMRI data while subjects were exposed to the sounds of human speech (vocal), and other non-vocal sounds. Each subject was exposed to 20 sounds of each type, totaling in n=40 trials in each scan. The study was rather large and consisted of about 200 subjects. The data was kindly made available by the authors at the OpenfMRI website<sup>3</sup>.

We perform group inference using within-subject permutations along the analysis pipeline of Stelzer et al. [2013], which was also reported in Gilron et al. [2016]. For completeness, the pipeline is described in Appendix A. To demonstrate our point, we compare the sd population test with the svm.cv.1 accuracy test.

In agreement with our simulation results, the population test (sd) discovers more brain regions of interest when compared to an accuracy test (svm.cv.1). The former discovers 1, 232 regions, while the latter only 441, as depicted in Figure 2. We emphasize that both test statistics were compared with the same permutation scheme, and the same error controls, so that any difference in detections is due to their different power.

<sup>3</sup>https://openfmri.org/

Having established that accuracy tests are typically underpowered for signal detection compared to population tests, we wish to identify the conditions under which this will occur, and discuss practical implications.

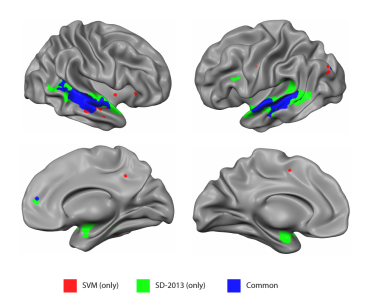


Figure 2: Brain regions encoding information discriminating between vocal and non-vocal stimuli. Map reports the centers of 27-voxel sized spherical regions, as discovered by an accuracy test (svm.cv.1), and a population test (sd). svm.cv.1 was computed using 5-fold cross validation, and a cost parameter of 1. Region-wise significance was determined using the permutation scheme of Stelzer et al. [2013], followed by region-wise  $FDR \leq 0.05$  control using the Benjamini-Hochberg procedure [Benjamini and Hochberg, 1995]. Number of permutations equals 400. The population test detect 1, 232 regions, and the accuracy test 441, 399 of which are common to both. For the details of the analysis see Appendix A and Gilron et al. [2016].

#### 6 Discussion

We have set out to understand which of the tests is more powerful: the accuracy test or the population test. No amount of simulations can replace the insight provided by a good closed-form analytic result. The finite sample power of permutation tests is a formidable mathematical problem, so we currently content ourselves with simulations We have concluded that the population tests are typically preferable. Their high dimensional versions, such as Srivastava [2007] and Schäfer and Strimmer [2005], are particularly well suited for neuroimaging problems such as MVPA. We attribute this

to several phenomena: (a) Discretization introduced in finite samples by the accuracy test statistic. (b) Inefficient use of the data for the validation holdout set. (c) Regularization crucial in high dimensional problems.

The presence of heavy tails shrinks the power advantage of the population tests over accuracy tests. Our empirical example suggests that even if the population test does not necessarily dominate the accuracy test in power, empirically, it does have an advantage.

The degree of discretization is governed by the sample size. For this reason, an asymptotic analysis such as Ramdas et al. [2016] may uncover the holdout inefficiency, but will not uncover the discretization effect.

The practical advice for the practitioner, is that for the purpose of signal detection, there is typically a population test that is more powerful than an accuracy test. There is also a good chance that it would be easier to implement, and faster to run, since no cross validation will be involved.

#### 6.1 Ease of implementation

A very important consideration is the ease of implementation. The need for cross validation of the accuracy test greatly increases its computational com-plexity. Moreover, anyone who has actually implemented tests with discrete statistics, will attest they are more prone to programming errors. This is because their unforgiveness to the type of inequalities used. Indeed, mistak-enly replacing a weak inequality with a strong inequality in one's program may considerably change the results. This is not the case for continuous test statistics. 

#### 6.2 Reservations

Some reservations to the generality of our findings are in order. Firstly, not all accuracy tests are concerned with signal detection. Consider brain decoding for machine interfaces, or clinical diagnosis, where the presence of a medical condition is predicted from imaging data [e.g. Olivetti et al., 2012, Wager et al., 2013]. In those examples, the purpose of the test is not to detect a difference between classes, but to actually test the performance of a particular classifier.

Secondly, it may be argued that accuracy tests permits the separation between classes in high dimensions, such as in *reproducing kernel Hilbert spaces* (RKHS) by using non-linear predictors. This is a false argument–accuracy test do not have any more flexibility that population tests. Indeed, it is possible to test for location in the same dimension the classifier is learned. Gretton et al. [2012] is an example where the test for location is performed

in the RKHS of the data. It is also possible to test for the equality of two multivariate distributions [TODO: cite vogelstein]. On the other hand, based on our reported neuroimaging example, and others, we find that a population test in the original feature space is indeed a simple and powerful approach to signal detection.

#### 249 6.3 A good accuracy test

For the cases a population test cannot replace an accuracy test, we collect some conclusions and best practices from our simulations. We give particular emphasis in this section to V-fold cross validation due to it popularity, but note that sampling the test set with replacement is actually preferable, as we discuss in Section 6.4.

Sample size. The conservativeness of accuracy tests decrease with sample size.

Permute features. Permuting features is easier than permuting labels.

It allows to preserve balanced folds after a permutation without refolding.

Although we not we did not find a power difference between balanced and unbalanced foldings.

Use less folds. For V-fold CV, power decreases as the number of folds increases. This is quite interesting since two phenomena compete as the number of folds increase: (a) the train set is larger so that better accuracies are achievable. (b) The test set is smaller so that the accuracy estimate is more variable. The decrease in power with increase fold number suggests that the latter dominates the former. Put differently: it is easier to detect a small stable departure from chance level, than a large but unstable one.

Resubstitution accuracy in low dimension. Resubstitution accuracy 268 useful in low dimension. In high dimension, the power loss is considerable 269 compared to a cross validated approach. We attribute this to the compound-270 ing of discretization and concentration effects: the difference between the 271 sampling distribution of the resubstitution accuracy is simply indistinguish-272 able under the null and under the alternative. In low dimensional problems, 273 the discretization is less impactful, and the computational burden of cross 274 validation can be avoided by using the resubstitution accuracy. There is 275 a fundamental difference between V-folding and resubstitution. The latter 276 should not be thought of as the limit of the former.

Regularize Regularizing the accuracy test proves very useful in high dimensional problems. Put differently: reducing variance by adding some bias is very useful to detect better-than-chance classification.

Don't z-score. There is no gain in z-scoring the accuracy scores. Our motivating rational was clearly flawed. [TODO: why?]

#### 6.4 Smoothing accuracy estimates

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It may be possible to alleviate the effect of discretization by appropriate 284 cross-validation. The discreteness of the accuracy statistic is governed by 285 the number of examples in the union (over all validation iterations) of test 286 sets. For V-fold CV, for instance, this number is simply the sample size. This 287 suggests that the accuracy can be "smoothed" by allowing the test sample to 288 be drawn with replacement. The bootstrap may seem like a good candidate 289 approach since it samples examples with replacement. It does so, however, for the train set, and not the test set. An algorithm that samples test sets with replacement is the leave-one-out bootstrap estimator (bLOO) and its 292 derivation—the 0.632 bootstrap estimator (b0.632) [Hastie et al., 2003, Sec 293 7.11]. 294

**Definition 1** (bLOO). Denoting by  $C^{(i)}$  the index set of bootstrap samples, b, where observation i is not in the train set, and by  $\mathcal{A}(\mathcal{S})^b$  the classifier fitted to the b'th bootstrap training sample, then the leave-one-out bootstrap estimate is defined as:

$$\mathcal{E}_{bLOO} := \frac{1}{n} \sum_{i=1}^{n} \frac{1}{|C^{(i)}|} \sum_{b \in C^{(i)}} I(\mathcal{A}(\mathcal{S})^{b}(x_{i}) = y_{i}).$$

Equivalently, denoting by  $S^{(b)}$  the indexes of observations, i, that are not in the bootstrap train sample b,

$$\mathcal{E}_{bLOO} = \frac{1}{B} \sum_{b=1}^{B} \frac{1}{|S^{(b)}|} \sum_{i \in S^{(b)}} I(\mathcal{A}(\mathcal{S})^b(x_i) = y_i).$$

**Definition 2** (b0.632). Denoting by  $\mathcal{E}_{resub}$  the resubstitution accuracy estimate, the b0.632 accuracy estimator,  $\mathcal{E}_{0.632}$ , is defined as

$$\mathcal{E}_{0.632} := 0.368 \ \mathcal{E}_{resub} + 0.632 \ \mathcal{E}_{bLOO}.$$

Simulation results reported in Figure 3 with naming conventions in Table 2. It can be seen that selecting test sets with replacement does increase the power, when compared to V-fold cross validation, but still falls short from the power of population tests. It can also be seen that power increases with the number of bootstrap replications, itself reducing the level of discretization. The type of bootstrap, bLOO versus b0.632, does not change the power.

Name	Basis	Type	В	Accuracy	Parameters
lda.Boot.1	LDA	b0.632	10	accuracy	_
lda.Boot.2	LDA	bLOO	10	accuracy	_
svm.Boot.1	SVM	b0.632	10	accuracy	cost=1e1
svm.Boot.2	SVM	bLOO	10	accuracy	cost=1e1
svm.Boot.3	SVM	b0.632	50	accuracy	cost=1e1
svm.Boot.4	SVM	bLOO	50	accuracy	cost=1e1

Table 2: The same as Table 1 for bootstraped accuracy estimates. bLOO and b0.632 are defined in definitions 1 and 2 respectively. B denotes the number of Bootstrap samples.

### 6.5 High dimensional classifiers

Inspecting Figure 1a (for instance), it can be seen that Hotelling's  $T^2$  test has similar power to accuracy tests. It should thus be argued that the real advantage of the population tests is due to their adaptation to high dimension by regularization (sd and Hotelling.shrink), and not only to discretization. To study this, we call upon several regularized classifiers, designed for high dimensional problems. In the spirit of the regularized covariance of Hotelling.shrink, we try an  $l_2$  regularized sym Friedman et al. [2010], and shrinkage based LDA [Pang et al., 2009, Ramey et al., 2016]. In the spirit of the diagonalized covariance of sd, we try a diagonalized LDA [Dudoit et al., 2002], which can be thought of a method intersecting Fisher's LDA and Naive Bayes.

Simulation results reported in Figure 4 with naming conventions in Table 3. It can be seen that regularizing a classifier in high dimension, just like a parameter test, improves power. It can also be seen that (regularized) parameter tests are still more powerful than (regularized) accuracy tests. This was to be expected, since we already saw in (e.g. Figure 1a) that the unregularized parameter test, *Hotelling*, is slightly more powerful than the regularized accuracy test, *svm.CV.1* for instance.

We can compound regularization in this section with the bootstrapping

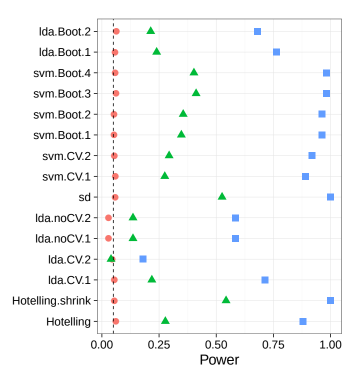


Figure 3: Bootstrap—The power of a permutation test with various test statistics. The power on the x axis. Effect are color and shape coded. The various statistics on the y axis. Their details are given in tables 1 and 2. Effects vary over 0 (red circle), 0.25 (green triangle), and 0.5 (blue square). Simulation details in Appendix B.

from Section 6.4, to improve finite sample power of the accuracy tests. This is done in the *svm.highdim.2* test, which still falls short from the power of the location tests, but is a much more powerful accuracy test than the original non-regularized, V-fold validated, version of *svm.CV.1*.

Name	Basis	CV	Accuracy	Parameters
svm.highdim.1	SVM	V-fold	accuracy	cost=1e-1
svm.highdim.2	SVM	B = 50	accuracy	cost=1e-1
lda.highdim.1	LDA	V-fold	accuracy	_
lda.highdim.2	LDA	V-fold	accuracy	_
lda.highdim.3	LDA	V-fold	accuracy	_

Table 3: The same as Table 1 for regularized (high dimensional) predictors. svm.highdim.1 is an l2 regularized SVM Friedman et al. [2010]. svm.highdim.2 is the same with b0.632 instead of V-fold cross validation. lda.highdim.1 is the Diagonal Linear Discriminant Analysis of Dudoit et al. [2002]. lda.highdim.2 is the High-Dimensional Regularized Discriminant Analysis of Ramey et al. [2016]. lda.highdim.3 is the Shrinkage-based Diagonal Linear Discriminant Analysis of Pang et al. [2009].



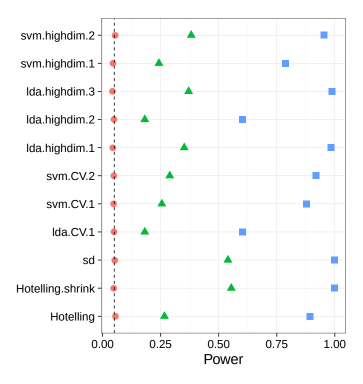


Figure 4: **HighDim Classifier**— The power of a permutation test with various test statistics. The power on the x axis. Effect are color and shape coded. The various statistics on the y axis. Their details are given in tables 1 and 3. Effects vary over 0 (red circle), 0.25 (green triangle), and 0.5 (blue square). Simulation details in Appendix B.

#### 6.6 Related Literature

Ojala and Garriga [2010] study the power of two accuracy tests: one testing the "no signal" null hypothesis, and the other testing the "independent features" null hypothesis. They perform an asymptotic analysis, and a simulation study. They also apply various classifiers to various data sets. Their emphasis is the effect of the underlying classifier on the power, and the potential of the "independent features" test for feature selection. This is a very different emphasis from our own.

Olivetti et al. [2012] and Olivetti et al. [2014] looked into the problem of choosing a good accuracy test. They propose a new test they call an *independence test*, and demonstrate by simulation that it has more power than other accuracy tests, and can deal with non-balanced data sets. We did not include this test in the battery we compared, but we note the following: (a) The independence test of Olivetti et al. [2012] relies on a discrete test statistic. It may thus be improved with the methods discussed in this section, before the application of Olivetti et al. [2012]'s independence test. (b) In contrast with the underlying motivation of Olivetti et al. [2012]'s independence test, we did not find that balancing the data folds is crucial for an accuracy test.

Golland et al. [2005] study accuracy tests using simulation, neuroimaging data, genetic data, and analytically. Their analytic results formalize our intuition from Section 1 on the effect of concentration of the accuracy statistic: The finite Vapnik–Chervonenkis (VC) dimension requirement [Golland and Fischl, 2003, Sec 4.3] prevents the permutation p-value from (asymptotically) concentrating near 1. Like ourselves, they also find that the power increases with the size of the test set (Figure 4, middle). This is seen in their Figure 4, where the size of the test-set, K, governs the discretization. Since they permute features, not labels, then all their permutation samples are balanced, and there is no issue of refolding.

Golland et al. [2005] simulate the power of accuracy tests by sampling from a Gaussian mixture family of models, and not from a location family as our own simulations. Under their model  $(x_i|y_i=1) \sim p\mathcal{N}(\mu_1,I) + (1-p)\mathcal{N}(\mu_2,I)$  and  $(x_i|y_i=-1) \sim (1-p)\mathcal{N}(\mu_1,I) + p\mathcal{N}(\mu_2,I)$ . Varying p interpolates between the null distribution (p=0.5) and a location shift model (p=0). We now perform the same simulation as Golland et al. [2005], after parameterizing p so that p=0 corresponds to the null model, and in the same dimensionality as our previous simulations We find that also in this mixture class of models a population test has more power than an accuracy test (Figure 5).

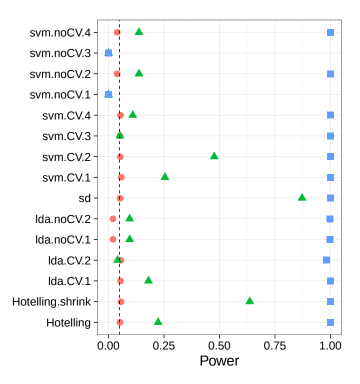


Figure 5: Mixture—  $\mathbf{x}_i = \chi_i \mu + \eta_i$ ;  $\chi_i = \{-1, 1\}$  and  $Prob(\chi_i = 1) = (1/2 - p)^{\mathbf{y}_i^*} (1/2 + p)^{1-\mathbf{y}_i^*}$ .  $\mu$  is a p-vector with  $3/\sqrt{p}$  in all coordinates. The effect, p, is color and shape coded and varies over 0 (red circle), 1/4 (green triangle) and 1/2 (blue square).

### 366 6.7 Epilogue

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Given all the above, we find the popularity of accuracy tests quite puzzling.
We believe this is due to a reversal of the inference cascade. Researchers first
fit a classifier, and then ask if the classes are any different. Were they to
start by asking if classes are any different, and only then try to classify, then
population tests would naturally arise as the preferred method. As put by
Ramdas et al. [2016]:

The recent popularity of machine learning has resulted in the extensive teaching and use of prediction in theoretical and applied communities and the relative lack of awareness or popularity of the topic of Neyman-Pearson style hypothesis testing in the computer science and related "data science" communities.

And more simply by Frank Harrell in the CrossValidated Q&A site<sup>4</sup>:

<sup>4</sup>http://stats.stackexchange.com/questions/17408/how-to-assess-statistical-significance-of-the-accuracy-of-a-classifier.

379 ... your use of proportion classified correctly as your accuracy 380 score. This is a discontinuous improper scoring rule that can be 381 easily manipulated because it is arbitrary and insensitive.

# 7 Acknowledgments

## 3 References

- T. W. Anderson. An Introduction to Multivariate Statistical Analysis. Wiley-Interscience, Hoboken, NJ, 3 edition edition, July 2003. ISBN 978-0-471-36091-9.
- Y. Benjamini and Y. Hochberg. Controlling the false discovery rate: a practical and powerful approach to multiple testing. *JOURNAL-ROYAL STA-TISTICAL SOCIETY SERIES B*, 57:289–289, 1995.
- S. Dudoit, J. Fridlyand, and T. P. Speed. Comparison of Discrimination
   Methods for the Classification of Tumors Using Gene Expression Data.
   Journal of the American Statistical Association, 97(457):77–87, Mar. 2002.
   ISSN 0162-1459. doi: 10.1198/016214502753479248.
- J. Friedman, T. Hastie, and R. Tibshirani. Regularization Paths for General ized Linear Models via Coordinate Descent. *Journal of Statistical Software*,
   33(1):1–22, 2010.
- R. Gilron, J. Rosenblatt, O. Koyejo, R. A. Poldrack, and R. Mukamel. Quantifying spatial pattern similarity in multivariate analysis using functional anisotropy. arXiv:1605.03482 [q-bio], May 2016.
- P. Golland and B. Fischl. Permutation tests for classification: towards statistical significance in image-based studies. In *IPMI*, volume 3, pages 330–341.
   Springer, 2003.
- P. Golland, F. Liang, S. Mukherjee, and D. Panchenko. Permutation Tests for Classification. In P. Auer and R. Meir, editors, *Learning Theory*, number 3559 in Lecture Notes in Computer Science, pages 501–515. Springer Berlin Heidelberg, June 2005. ISBN 978-3-540-26556-6 978-3-540-31892-7. doi: 10.1007/11503415\_34.
- T. R. Golub, D. K. Slonim, P. Tamayo, C. Huard, M. Gaasenbeek, J. P. Mesirov, H. Coller, M. L. Loh, J. R. Downing, M. A. Caligiuri, C. D. Bloomfield, and E. S. Lander. Molecular Classification of Cancer: Class Discovery and Class Prediction by Gene Expression Monitoring. Science, 286(5439):531–537, Oct. 1999. ISSN 0036-8075, 1095-9203. doi: 10.1126/science.286.5439.531.
- A. Gretton, K. M. Borgwardt, M. J. Rasch, B. Schölkopf, and A. Smola. A
   Kernel Two-sample Test. J. Mach. Learn. Res., 13:723–773, Mar. 2012.
   ISSN 1532-4435.

- T. Hastie, R. Tibshirani, and J. Friedman. *The Elements of Statistical Learning*. Springer, July 2003. ISBN 0-387-95284-5.
- J. Hemerik and J. Goeman. Exact testing with random permutations. arXiv:1411.7565 [math, stat], Nov. 2014.
- H. Hotelling. The Generalization of Student's Ratio. *The Annals of Mathematical Statistics*, 2(3):360–378, Aug. 1931. ISSN 0003-4851, 2168-8990. doi: 10.1214/aoms/1177732979.
- W. Jiang, S. Varma, and R. Simon. Calculating confidence intervals for
   prediction error in microarray classification using resampling. Statistical
   Applications in Genetics and Molecular Biology, 7(1), 2008.
- L. Juan and H. Iba. Prediction of tumor outcome based on gene expression data. Wuhan University Journal of Natural Sciences, 9(2):177–182, Mar. 2004. ISSN 1007-1202, 1993-4998. doi: 10.1007/BF02830598.
- N. Kriegeskorte, R. Goebel, and P. Bandettini. Information-based functional brain mapping. *Proceedings of the National Academy of Sciences of the* United States of America, 103(10):3863–3868, July 2006. ISSN 0027-8424, 1091-6490. doi: 10.1073/pnas.0600244103.
- E. L. Lehmann. Parametric versus nonparametrics: two alternative methodologies. *Journal of Nonparametric Statistics*, 21(4):397–405, 2009. ISSN 1048-5252. doi: 10.1080/10485250902842727.
- G. J. McLachlan. The bias of the apparent error rate in discriminant analysis.
   Biometrika, 63(2):239-244, Jan. 1976. ISSN 0006-3444, 1464-3510. doi: 10.1093/biomet/63.2.239.
- S. Mukherjee, P. Tamayo, S. Rogers, R. Rifkin, A. Engle, C. Campbell,
   T. R. Golub, and J. P. Mesirov. Estimating dataset size requirements
   for classifying DNA microarray data. *Journal of Computational Biology:* A Journal of Computational Molecular Cell Biology, 10(2):119–142, 2003.
   ISSN 1066-5277. doi: 10.1089/106652703321825928.
- M. Ojala and G. C. Garriga. Permutation Tests for Studying Classifier Performance. Journal of Machine Learning Research, 11(Jun):1833–1863, 2010.
   ISSN ISSN 1533-7928.
- E. Olivetti, S. Greiner, and P. Avesani. Induction in Neuroscience with Classification: Issues and Solutions. In G. Langs, I. Rish, M. Grosse-Wentrup, and B. Murphy, editors, *Machine Learning and Interpretation*

- in Neuroimaging, number 7263 in Lecture Notes in Computer Science, 451
- pages 42–50. Springer Berlin Heidelberg, 2012. ISBN 978-3-642-34712-2 452
- 978-3-642-34713-9. doi: 10.1007/978-3-642-34713-9\_6. 453
- E. Olivetti, S. Greiner, and P. Avesani. Statistical independence for the 454 evaluation of classifier-based diagnosis. Brain Informatics, 2(1):13–19, Dec. 455
- 2014. ISSN 2198-4018, 2198-4026. doi: 10.1007/s40708-014-0007-6. 456
- H. Pang, T. Tong, and H. Zhao. Shrinkage-based Diagonal Discriminant 457
- Analysis and Its Applications in High-Dimensional Data. Biometrics, 65 458
- (4):1021–1029, Dec. 2009. ISSN 1541-0420. doi: 10.1111/j.1541-0420.2009. 459
- 01200.x.460
- F. Pereira, T. Mitchell, and M. Botvinick. Machine learning classifiers and 461
- fMRI: A tutorial overview. NeuroImage, 45(1, Supplement 1):S199-S209, 462
- Mar. 2009. ISSN 1053-8119. doi: 10.1016/j.neuroimage.2008.11.007. 463
- C. R. Pernet, P. McAleer, M. Latinus, K. J. Gorgolewski, I. Charest, P. E. G. 464
- Bestelmeyer, R. H. Watson, D. Fleming, F. Crabbe, M. Valdes-Sosa, and 465
- P. Belin. The human voice areas: Spatial organization and inter-individual 466
- variability in temporal and extra-temporal cortices. NeuroImage, 119:164– 467
- 174, Oct. 2015. ISSN 1053-8119. doi: 10.1016/j.neuroimage.2015.06.050. 468
- M. D. Radmacher, L. M. McShane, and R. Simon. A Paradigm for 469
- Class Prediction Using Gene Expression Profiles. Journal of Computa-470
- tional Biology, 9(3):505-511, June 2002. ISSN 1066-5277. doi: 10.1089/ 471
- 106652702760138592.472
- A. Ramdas, A. Singh, and L. Wasserman. Classification Accuracy as a Proxy
- for Two Sample Testing. arXiv:1602.02210 [cs, math, stat], Feb. 2016. 474
- J. A. Ramey, C. K. Stein, P. D. Young, and D. M. Young. High-Dimensional 475
- Regularized Discriminant Analysis. arXiv preprint arXiv:1602.01182,
- 2016. 477
- J. Schäfer and K. Strimmer. A Shrinkage Approach to Large-Scale Covariance 478
- Matrix Estimation and Implications for Functional Genomics. Statistical 479
- Applications in Genetics and Molecular Biology, 4(1), Jan. 2005. ISSN 480
- 1544-6115. doi: 10.2202/1544-6115.1175. 481
- D. K. Slonim, P. Tamayo, J. P. Mesirov, T. R. Golub, and E. S. Lander. Class 482
- Prediction and Discovery Using Gene Expression Data. In *Proceedings of* 483
- the Fourth Annual International Conference on Computational Molecular 484

- Biology, RECOMB '00, pages 263–272, New York, NY, USA, 2000. ACM.
   ISBN 978-1-58113-186-4. doi: 10.1145/332306.332564.
- M. S. Srivastava. Multivariate Theory for Analyzing High Dimensional Data.
   Journal of the Japan Statistical Society, 37(1):53–86, 2007. doi: 10.14490/
   jjss.37.53.
- M. S. Srivastava, S. Katayama, and Y. Kano. A two sample test in high
   dimensional data. *Journal of Multivariate Analysis*, 114:349–358, Feb.
   2013. ISSN 0047-259X. doi: 10.1016/j.jmva.2012.08.014.
- J. Stelzer, Y. Chen, and R. Turner. Statistical inference and multiple testing correction in classification-based multi-voxel pattern analysis (MVPA): Random permutations and cluster size control. *NeuroImage*, 65:69–82, Jan. 2013. ISSN 1053-8119. doi: 10.1016/j.neuroimage.2012.09.063.
- A. W. van der Vaart. Asymptotic Statistics. Cambridge University Press, Cambridge, UK; New York, NY, USA, Oct. 1998. ISBN 978-0-521-49603-2.
- G. Varoquaux, P. R. Raamana, D. Engemann, A. Hoyos-Idrobo, Y. Schwartz,
   and B. Thirion. Assessing and tuning brain decoders: cross-validation,
   caveats, and guidelines. working paper or preprint, June 2016.
- T. D. Wager, L. Y. Atlas, M. A. Lindquist, M. Roy, C.-W. Woo, and E. Kross.
   An fMRI-Based Neurologic Signature of Physical Pain. New England Jour nal of Medicine, 368(15):1388–1397, Apr. 2013. ISSN 0028-4793. doi: 10.1056/NEJMoa1204471.

## A Analysis pipeline

Here is the analysis pipeline of Stelzer et al. [2013] we for the auditory data in Gilron et al. [2016]. Denoting by  $i=1,\ldots,I$  the subject index,  $v=1,\ldots,V$  the voxel index, and  $s=1,\ldots,S$  the permutation index. Since regions<sup>5</sup> are centered around a unique voxel, the voxel index v also serves as a unique region index. Algorithm 1 computes a region-wise test statistic, which is compared to its permutation null distribution computed by Algorithm 2.

#### Algorithm 1: Compute a group parametric map.

Data: fMRI scans, and experimental design.

**Result:** Brain map of group statistics:  $\{\bar{T}_v\}_{v=1}^V$ 

1 for  $v \in 1, \ldots, V$  do

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for  $i \in 1, \ldots, I$  do

**3**  $T_{i,v} \leftarrow \text{test statistic for subject } i \text{ in a region centered at } v.$ 

4  $\bar{T}_v \leftarrow \frac{1}{I} \sum_{i=1}^{I} T_{i,v}$ .

#### Algorithm 2: Compute a permutation p-value map.

Data: fMRI scans of 20 subjects, experimental design.

**Result:** Brain map of permutation p-values:  $\{p_v\}_{v=1}^V$ 

1 for  $s \in 1, \dots S$  do

2 permute labels;

 $\mathbf{3} \quad | \quad \bar{T}_v^s \leftarrow \text{parametric map}$ 

 $<sup>^5</sup>searchlight$  or sphere in the MVPA parlance

## B Simulation Details

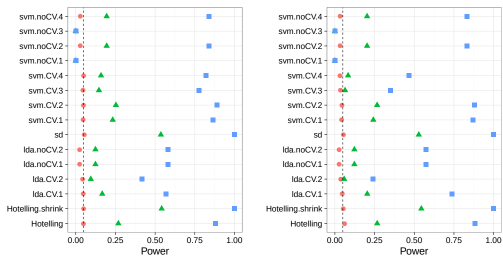
The following details are common to all the reported simulations, unless stated otherwise in a figure's caption. The R code for the simulations can be found in [TODO].

Each simulation is based on 4,000 replications. In each replication, we generate n i.i.d. samples from a shift model  $\mathbf{x}_i = \mu \mathbf{y}_i^* + \eta_i$ . Where  $y_i^* = \{0, 1\}$  is the class of subject i in dummy coding. Recalling that  $y_i = \{-1, 1\}$  is the class in effect coding, then clearly  $y_i = 2y_i^* - 1$ . The noise is distributed as  $\eta_i \sim \mathcal{N}_p(0, \Sigma)$ . The sample size n = 40. The dimension of the data is p = 23. The covariance  $\Sigma = I$ . Effects, i.e. shifts  $\mu$ , are equal coordinate p-vectors with coordinates that vary over  $\mu \in \{0, 1/4, 1/2\}$ .

Having generated the data, we compute each of the test statistics in Table 1. For test statistics that require data folding, we used 8 folds. We then compute a permutation p-value by permuting the class labels, and recomputing each test statistic. We perform 400 such permutations. We then reject the  $\mu_i=0$  null hypothesis if the permutation p-value is smaller than 0.05. The reported power is the proportion of replication where the permutation p-value falls below 0.05.

# 534 C Simulation Results

 $Figure \ 6$ : Simulation details in Appendix B except the changes in the sub-captions.



- (a) 2-fold cross validation. Balanced folding.
- (b) 20-fold cross validation. Balanced folding

Figure 7: Simulation details in Appendix B except the changes in the sub-captions.



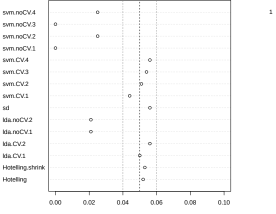
- (a) 2-fold cross validation. Unbalanced folding.
- (b) 20-fold cross validation.
  Unbalanced folding.

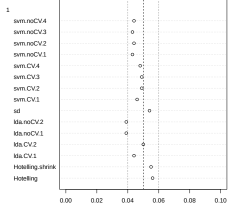
Figure 8: Simulation details in Appendix B except the changes in the sub-captions.



- (a) Scale Change—  $\mathbf{x}_i = \eta_i * \mu^{\mathbf{y}_i^*}$  so that the effect are a scale change.
- (b) Heavytailed- $\eta_i$  is not p-variate Gaussian, but rather p-variate t, with df = 3.

Figure 9: Simulation details in Appendix B except the changes in the sub-captions.

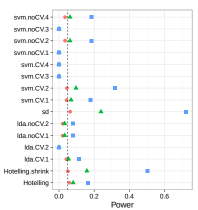




- (a) Low-Dimension—False positive rates for n = 40.
- (b) High-Dimension—False positive rates for n = 400.

Figure 10: Simulation details in Appendix B except the changes in the sub-captions.





(a) High-Dimension, local alternative—  $n=400, \\ \mu \in \frac{1}{\sqrt{10}} \times \{0,1/4,1/2\}.$ 

(b) AR(1) dependence—  $\Sigma_{k,l} = \rho^{|k-l|}; \rho = 0.8.$