### AutoML: Evaluation

Benchmarking and Comparing Learners

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### Benchmark Experiments

- different learning algorithms applied to one or more data sets to compare and rank their performances
- synchronized train and test sets, i.e. the same resampling method with the same train-test splits should be used to determine performance
   Example: Benchmark results (per CV-fold) of CART and random forest using 2-fold CV with MSE as performance measure:

data set	k-th fold	MSE (rpart)	MSE (randomForest)
BostonHousing	1	29.4	17.13
BostonHousing	2	20.5	8.90
mtcars	1	35.0	7.53
mtcars	2	38.9	6.73

# Hypothesis Testing in Benchmarking I

We want to know if the difference in performance between models (or algorithms) is significant or only by chance.

### Null Hypothesis Statistical Testing (NHST) in Benchmarking:

- ullet formulate null hypothesis  $H_0$  (e.g. the expected generalization error of two algorithms is equivalent) and alternative hypothesis  $H_1$
- ullet use hypothesis test to reject  $H_0$  (not rejecting  $H_0$  does not mean that we accept it)
- ullet rejecting  $H_0$  gives some confidence that the observed results may not be random

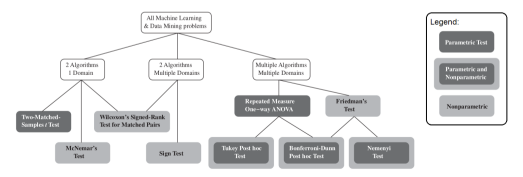
#### Typical example in machine learning:

- $H_0$ : on average, model 1 does not perform better than model 2
- $\bullet$   $H_1$ : on average, model 1 outperforms model 2
- Aim: Reject  $H_0$  with confidence level of  $1-\alpha$  (common values for  $\alpha$  include 0.05 and 0.01)

## Hypothesis Testing in Benchmarking II

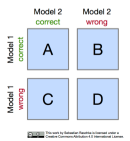
Selection of an appropriate hypothesis test is at least based on the type of problem, i.e. if the aim is to compare

- 2 models / algorithms on a single domain (i.e. on a single data set)
- 2 algorithms across different domains (i.e. on multiple data sets)
- multiple algorithms across different domains / data sets



#### McNemar Test I

- non-parametric test used on paired dichotomous nominal data; does not make any distributional assumptions beyond statistical independence of samples
- pairs are e.g. labels predicted by different models on the same data
- compares the classification accuracy of two models
- both models trained and evaluated on the exact same training and test set; contingency table based on two paired vectors that indicate whether each model predicted an observation correctly



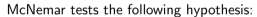
- A: #obs. correctly classified by both
- B: #obs. only correctly classified by model 1
- C: #obs. only correctly classified by model 2
- D: #obs. misclassified by both

#### McNemar Test II

Error of each model can be computed as follows:

- Model 1: (C+D)/(A+B+C+D)
- Model 2: (B+D)/(A+B+C+D)

Even if the models have the **same** errors (indicating equal performance), cells B and C may be different because the models may misclassify different instances.

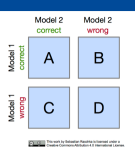


- ullet  $H_0$ : both models have the same performance (we expect  $\mathsf{B}=\mathsf{C}$ )
- ullet  $H_1$ : performances of the two models are not equal

The test statistic is computed as

$$\chi^2_{Mc} = \frac{(|B-C|-1)^2}{B+C} \sim \chi^2_1.$$

**Note**: The McNemar test should only be used if B + C > 20.



### McNemar Test III

#### Example:

$$\begin{tabular}{c|ccccc} & & Random Forest \\ \hline & & 0 & 1 \\ \hline \\ Tree & 0 & 30 & 5 \\ 1 & 17 & 42 \\ \hline \end{tabular}$$

Calculating the test statistic:

$$\chi_{Mc}^2 = \frac{(|5-17|-1)^2}{5+17} = 5.5 > 3.841 = \chi_{1,0.95}^2$$

We can reject  $H_0$  at a significance level of 0.05, i.e. we reject the hypothesis that the tree and the random forest have the same performance.

Significance level must be chosen before applying the test (avoid p-value hacking).

### Two-Matched-Samples t-Test I

- two-matched-samples t-test (i.e. a paired t-test) is the simplest hypothesis test to compare two **models** on a single test set based on arbitrary performance measures
- parametric test and distributional assumptions must be made (which are often problematic):

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(pseudo-)normality usually met when sample size > 30 i.i.d. samples usually met if loss of individual observations from single test set considered equal variances of populations can be investigated through plots
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### Two-Matched-Samples t-Test II

Compare two different models  $\hat{f}_1$  and  $\hat{f}_2$  w.r.t. performance measure calculated on test set of size  $n_{\text{test}}$ :

- $H_0$ :  $GE(\hat{f}_1) = GE(\hat{f}_2)$  vs.  $H_1$ :  $GE(\hat{f}_1) \neq GE(\hat{f}_2)$
- test statistic  $T = \sqrt{n_{\text{test}}} \frac{\bar{d}}{\sigma_d}$  where
  - lacktriangle mean performance difference of both models is  $ar{d} = \hat{GE}_{\mathcal{D}_{\mathsf{test}}}(\hat{f}_1) \hat{GE}_{\mathcal{D}_{\mathsf{test}}}(\hat{f}_2)$
  - standard deviation of this mean difference is

$$\sigma_d = \sqrt{\frac{1}{n_{\mathsf{test}} - 1} \sum_{i=1}^{n_{\mathsf{test}}} \left(d_i - \bar{d}\right)^2},$$

where 
$$d_i = L(y^{(i)}, \hat{f}_1(\mathbf{x}^{(i)})) - L(y^{(i)}, \hat{f}_2(\mathbf{x}^{(i)}))$$
 and  $\bar{d} = \frac{1}{n_{\text{test}}} \sum_{i=1}^{n_{\text{test}}} d_i$ 

**Note**:  $d_i$  is the difference of the outer loss of individual observations from the test set between the two models to be compared.

### Two-Matched-Samples t-Test III

- could also use a k-fold CV paired t-test to compare two algorithms (instead of two models) on single data set
- instead of comparing outer loss of individual observations, compare generalization errors per CV fold (i.e. k generalization errors for k CV folds)
- performance differences are not independent across CV folds due to overlapping training sets (which violates the assumption of i.i.d. samples)
- to partly overcome issue of overlapping training sets across folds, Dietterich suggests using 5 times 2-fold CV so that at least within each repetition neither training nor test sets overlap [Dietterich. 1998]

#### Friedman Test I

Compare multiple classifiers on multiple data sets:

- ullet  $H_0$ : all algorithms are equivalent in their performance and hence their average ranks should be equal
- ullet  $H_1$ : the average ranks for at least one algorithm is different

To evaluate n data sets and k algorithms:

- rank each algorithm on each data set from best-performing algorithm (rank 1) to worst-performing algorithm using any performance measure
- ullet  $R_{ij}$  is the rank of algorithm j on data set i
- if there is a d-way tie after rank r, assign rank of  $\left[(r+1)+(r+2)+...+(r+d)\right]/d$  to each tied classifier

#### Friedman Test II

#### Can now compute:

- overall mean rank  $ar{R} = rac{1}{nk} \sum_{i=1}^n \sum_{j=1}^k R_{ij}$
- sum of squares total  $SS_{Total}=n\sum_{j=1}^k(\bar{R}_{.j}-\bar{R})^2$  where  $\bar{R}_{.j}=\frac{1}{n}\sum_{i=1}^nR_{ij}$
- sum of squares error  $SS_{Error} = \frac{1}{n(k-1)} \sum_{i=1}^{n} \sum_{j=1}^{k} (R_{ij} \bar{R})^2$

Test statistic calculated as:

$${\chi_F}^2 = rac{SS_{Total}}{SS_{Error}} \sim \chi^2_{k-1}$$
 for large n (>15) and k (>4)

For smaller n and k, the  $\chi^2$  approximation is imprecise and a look up of  $\chi^2_F$  values that were approximated specifically for the Friedman test is suggested.

#### Post-Hoc Tests I

- Friedman test checks if all algorithms are ranked equally
- does not allow to identify best-performing algorithm
- $\rightarrow \text{ post-hoc tests}$

#### Post-hoc Nemenyi test:

- ullet compares all pairs of algorithms to find best-performing algorithm after  $H_0$  of the Friedman-test was rejected
- ullet for n data sets and k algorithms,  $\frac{k(k-1)}{2}$  comparisons
- calculate average rank of algorithm j on all n data sets:  $\bar{R}_{.j} = \frac{1}{n} \sum_{i=1}^{n} R_{ij}$

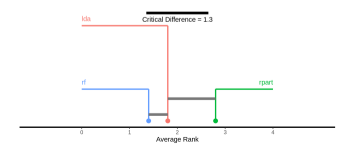
For any two algorithms  $j_1$  and  $j_2$ , test statistic computed as:

$$q = \frac{\bar{R}_{.j_1} - \bar{R}_{.j_2}}{\sqrt{\frac{k(k+1)}{6n}}}$$

#### Post-Hoc Tests II

#### Critical Difference Plot:

- quick way to see what differences are significant across all compared learners
- all learners that do not differ by at least the critical difference are connected by line
- a learner not connected to another learner and of lower rank can be considered better according to the chosen significance level



#### Post-Hoc Tests III

#### Post-hoc Bonferonni-Dunn test:

- ullet compares all algorithms with baseline (i.e. k-1 comparisons)
- used after Friedman test to find which algorithms differ from the baseline significantly
- uses Bonferonni correction to prevent randomly accepting one of the algorithms as significant due to multiple testing

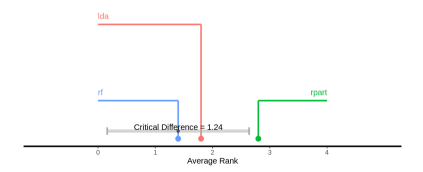
The test statistic is the same as before:

$$q = \frac{\bar{R}_{.j_1} - \bar{R}_{.j_2}}{\sqrt{\frac{k(k+1)}{6n}}}.$$

The performance of  $j_1$  and  $j_2$  are significantly different when  $|q| > q_{\alpha}$ , where the critical value  $q_{\alpha}$  is obtained from a table of the studentized range statistic divided by  $\sqrt{2}$ .

#### Post-Hoc Tests IV

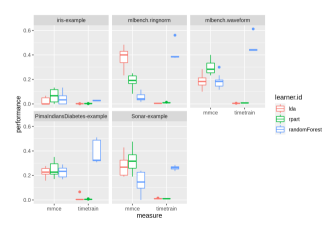
• learners within the baseline interval (gray line) perform not significantly different from the baseline



# Comparing Visually I

It can be helpful to inspect distributions visually for additional insights, e.g.

### **Boxplots**



# Comparing Visually II

### Rank plots

