Generalisation in theory and practice

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Classification

The classifier as a decision rule

A decision rule $\pi(a|x)$ generates a decision $a \in [m]$. It is the conditional probability of a given x.

A note on conditional probabilities

Even though normally conditional probabilities are defined as $P(A|B) = P(A \cap B)/P(B)$, the probability of the decision a is undefined without a given x. So it's better to think if $\pi(a|x)$ as a collection of distributions on a, one for each value of x.

Deterministic predictions given a model P(y|x)

Here, we pick the most likely class:

$$\pi(a|x_t) = \mathbb{I}\left\{a = \argmax_{y} P(y|x_t)\right\}$$

Deterministic predictions given a model P(y|x)

Here, we randomly select a class according to our model:

$$\pi(a|x_t)=P(y_t=a|x_t)$$

Accuracy as a classification metric

The accuracy of a single decision

$$U(a_t, y_t) = \mathbb{I}\left\{a_t = y_t\right\} = \begin{cases} 1, & \text{if } a_t = y_t \\ 0, & \text{otherwise} \end{cases}$$

The accuracy on the training set

$$U(\pi, D) \triangleq \frac{1}{T} \sum_{t=1}^{T} \sum_{a=1}^{m} \pi(y_t|x_t)$$

The expected accuracy of a decision rule

If $(x, y) \sim P$, the accuracy U of a stochastic decision rule π under the distribution P is the probability it predicts correctly

$$U(\pi, P) \triangleq \int_X dP(x) \sum_{y=1}^m P(y|x) \pi(y|x)$$

Regression

The regressor as a decision rule

A decision rule π generates a decision $a \in \mathbb{R}^m$.

- For randomised rules, $\pi(a|x)$ is the conditional density of a given x.
- ▶ For deterministic rules $\pi(x)$ is the prediction for x.

Mean-Squared Error on a Dataset

The mean-square error is simply the squared difference in predicted versus actual values:

$$\frac{1}{T} \sum_{t=1}^{T} [y_t - \pi(x_t)]^2$$

Expected MSE

If $(x,y)\sim P$, the expected MSE of a deterministic decision rule $\pi:X\to\mathbb{R}$ is

$$\int_X \int_Y dP(x,y)[y-\pi(x)]^2.$$

Training and overfitting

Training data

- \triangleright $D = ((x_t, y_t) : t = 1, ..., T).$
- \triangleright $x_t \in X, y_t \in Y.$

Assumption: The data is generated i.i.d.

- \blacktriangleright $(x_t, y_t) \sim P$ for all t (identical)
- $ightharpoonup D \sim P^T$ (independent)

The optimal decision rule for P

$$\max_{\pi} U(\pi, P) = \max_{\pi} \int_{X \times Y} dP(x, y) \sum_{a} \pi(a|x) U(a, y)$$

The optimal decision rule for D

$$\max_{\pi} U(\pi, D) = \max_{\pi} \sum_{(x,y) \in D} \sum_{a} \pi(a|x)U(a,y)$$

The Train/Validation/Test methodology

Main idea

Use each piece of data once to make decisions and measure

Training set

Use to decide low-level model parameters

Validation set

Use to decide between:

- ▶ different hyperparameters (e.g. K in nearest neighbours)
- model (e.g. neural networks versus kNN)

Test set

Use to measure the final quality of a model

Cross-validation (XV)

Idea

▶ Use XV to select hyperparameters instead of a single train/valid test.

Methodology

- ► Split training set *D* in *k* different subsets
- ► At iteration *i*
- Use the \$i\$-th subset for validation
- ▶ Use all the remaining k-1 subsets for training
- Average results on validation sets

Bootstrapping

Idea

- ► How to take into account variability?
- Resample the data and repeat your calculations for each resample

Boostrap samples

- ▶ Input: Data D, of size T
- For t in $\{1, \ldots, T\}$
- Select i uniformly in [T] Add the \$i\$-th point to D_b
 - ightharpoonup Return D_b

The wrong way to do XV for subset selection

- 1. Screen the predictors: find a subset of "good" predictors that show fairly strong (univariate) correlation with the class labels.
- 2. Using just this subset of predictors, build a multivariate classifier.
- 3. Use cross-validation to estimate the unknown tuning parameters and to estimate the prediction error of the final model.

Is this a correct application of cross-validation?

Consider a scenario with N=50 samples in two equal-sized classes, and p=5000 quantitative predictors (standard Gaussian) that are independent of the class labels. The true (test) error rate of any classifier is 50%.

The right way to do XV for feature selection

- 1. Divide the samples into K cross-validation folds (groups) at random.
- 2. For each fold $k = 1, 2, \ldots, K$
- Find a subset of "good" predictors that show fairly strong (univariate) correlation with the class labels, using all of the samples except those in fold k.
- 4. Using just this subset of predictors, build a multivariate classifier, using all of the samples except those in fold k.
- 5. Use the classifier to predict the class labels for the samples in fold k.

Learning and generalisation

How well can decision rule perform?

Estimation theory view

- Bias: The expected difference between the estimated value and the unknown parameter
- Variance: The expected difference between the estimated value and the unknown parameter

Learning theory view

- Approximation ability: How well a class of rules can approximate the optimal one.
- Statistical error: How easy it is to choose the best rule in the class.

The bias/variance trade-off

- ▶ Dataset $D \sim P$.
- ▶ Predictor $f_D(x)$
- ▶ Target function $y = f(x) + \epsilon$
- $ightharpoonup \mathbb{E}\,\epsilon = 0$ zero-mean noise with variance $\sigma^2 = \mathbb{V}(\epsilon)$

MSE decomposition

$$\mathbb{E}[(f - f_D)^2] = \mathbb{V}(f_D) + \mathbb{B}(f_D)^2 + \sigma^2$$

Variance

How sensitive the estimator is to the data

$$\mathbb{V}(f_D) = \mathbb{E}[(f_D - \mathbb{E}(f_D))^2]$$

Bias

What is the expected deviation from the true function

$$\mathbb{B}(f_D) \triangleq \mathbb{E}[(f_D - f)]$$

Example: mean estimation

- ightharpoonup Data $D=y_1,\ldots,y_T$ with $\mathbb{E}[y_t]=\mu$.
- ▶ Goal: estimate μ with some estimator f_D to minimise
- ► MSE: $\mathbb{E}[(y f_D)^2]$, the expected square difference between new samples our guess.

Optimal estimate

To minimise the MSE, we use $f^* = \mu$. This gives us two ideas:

Empirical mean estimator:

- $f_D = \sum_{t=1}^T x_t / T.$
- \blacktriangleright $\mathbb{V}(f_D) = \mathbb{E}[f_D \mu] = 1/\sqrt{T}$
- $ightharpoonup \mathbb{B}(f_D) = 0.$

Laplace mean estimator:

- $f_D = \sum_{t=1}^{T} (\lambda + x_t)/T.$
- $\blacktriangleright \mathbb{V}(f_D) = \mathbb{E}[f_D \mu] = \frac{1}{1 + \sqrt{T}}$
- $ightharpoonup \mathbb{B}(f_D) = O(1/T).$



A proof of the bias/variance trade-off

- $ightharpoonup RV's \ y_t \sim P, \ \mathbb{E}[y_t] = \mu, \ y_t = \mu + \epsilon_t.$
- ightharpoonup Estimator f_D , $D = y_1, \dots, y_{t-1}$.

$$\mathbb{E}[(f_{D} - y_{t})^{2}] = \mathbb{E}[f_{D}^{2}] - 2 \mathbb{E}[f_{D}y_{t}] + \mathbb{E}[y_{t}^{2}]$$

$$= \mathbb{V}[f_{D}] + \mathbb{E}[f_{D}]^{2} - 2 \mathbb{E}[f_{D}y_{t}] + \mathbb{E}[y_{t}^{2}]$$

$$= \mathbb{V}[f_{D}] + \mathbb{E}[f_{D}]^{2} - 2 \mathbb{E}[f_{D}] \mathbb{E}[y_{t}] + \mathbb{E}[y_{t}^{2}]$$

$$= \mathbb{V}[f_{D}] + \mathbb{E}[f_{D}]^{2} - 2 \mathbb{E}[f_{D}]\mu + \mathbb{E}[y_{t}^{2}]$$

$$= \mathbb{V}[f_{D}] + \mathbb{E}[f_{D}]^{2} - 2 \mathbb{E}[f_{D}]\mu + \mathbb{E}[(\mu + \epsilon_{t})^{2}]$$

$$= \mathbb{V}[f_{D}] + \mathbb{E}[f_{D}]^{2} - 2 \mathbb{E}[f_{D}]\mu + \mathbb{E}[\mu^{2} + 2\mu\epsilon_{t} + \epsilon_{t}^{2}]$$

$$= \mathbb{V}[f_{D}] + \mathbb{E}[f_{D}]^{2} - 2 \mathbb{E}[f_{D}]\mu + \mu^{2} + \sigma^{2}$$

$$= \mathbb{V}[f_{D}] + (\mathbb{E}[f_{D}] - \mu)^{2} + \sigma^{2}$$

$$= \mathbb{V}(f_{D}) + \mathbb{E}(f_{D})^{2} + \sigma^{2}$$

Generalisation error

Regret decomposition

Let the optimal rule be $\pi^* \in \Pi$, the best approximate rule be $\hat{\pi}^* \in \Pi$ and our rule be $\hat{\pi} \in \hat{\Pi}$. We call the difference between the performance of π^* and $\hat{\pi}$ our regret:

$$\underbrace{U(\pi^*,P)-U(\hat{\pi},P)}_{\text{regret}} = \underbrace{U(\pi^*,P)-U(\hat{\pi}^*,P)}_{\text{approximation error}} + \underbrace{U(\hat{\pi}^*,P)-U(\hat{\pi},P)}_{\text{estimation error}}$$

We can bound the regret by bounding each term separately.

- The approximation error tells us how expressive our class of rules is, i.e. how much we lose by looking at a restricted class $\hat{\Pi}$ of rules. It is similar to estimator bias.
- ► The statistical error tells us how well the empirical performance on *D* approximates the true performance. It is similar to estimator variance.
- As a rule of thumb, the larger our class, the better the possible approximation but the higher the statistical error.

Approximation error

- lackbox Our model limits us to a set of decision rules $\hat{\Pi}\subset \Pi$.
- ightharpoonup The most we could do is find the best rule in $\hat{\Pi}$.
- ► This still leaves a gap:

$$\Delta \triangleq \max_{\pi \in \Pi} U(\pi, P) - \max_{\hat{\pi} \in \hat{\Pi}} U(\pi, P)$$

The gap can be characterised in some cases.

Example: ϵ -net on Lipschitz $U(\cdot, P)$.

- Assume $U(\pi, P)$ is a Lipschitz function of π for all P, i.e. $|U(\pi, P) U(\pi', P)| \le Ld(\pi, \pi')$ for some metric d.
- ▶ Let $\hat{\Pi}$ be an ϵ -net on Π , i.e. $\max_{\pi \in \Pi} \min_{\pi' \in \hat{\Pi}} d(\pi, \pi') = \epsilon$.
- ▶ Then $\Delta \leq L\epsilon$.

Estimation error

- First, let us bound $U(\hat{\pi}^*, P) U(\hat{\pi}, P)$ by making an assumption.
- Then, we can prove that our assumption holds with high probability.

Lemma

Let $f,g:S\to\mathbb{R}$. If $\|f-g\|_\infty\leq\epsilon$ and $f(x)\geq f(z)$, while $g(y)\geq g(z)$, for all z, i.e. x,y maximise f,g respectively

$$f(x) - f(y) \le 2\epsilon$$
.

This holds as: $f(x) - f(y) \le g(x) + \epsilon - f(y) \le g(y) + \epsilon - f(y) \le 2\epsilon$.

Corollary

If $|U(\pi, P) - U(\pi, D)| \le \epsilon$ for all π then

$$U(\hat{\pi}^*, P) - U(\hat{\pi}, P) \le 2\epsilon$$

Let us now prove that, with high probability, $|U(\pi, P) - U(\pi, D)| \le \epsilon$.



Bounding the estimation error

For any fixed rule $\pi \in \Pi$ and utility function $U : \Pi \times X^T \to [0,1]$,

$$P^{T}(|U(\pi,D)-U(\pi,P)| \geq \epsilon) \leq 2 \exp(-2T\epsilon^{2}).$$

This is a direct application of Hoeffding's inequality¹. Taking the union bound over the set \hat{H} gives:

$$P^{T}(\exists \pi \in \hat{\Pi} : |U(\pi, D) - U(\pi, P)| \ge \epsilon) \le 2|\hat{\Pi}| \exp(-2T\epsilon^{2}).$$

Setting the right side equal to δ and re-arranging,

$$P^T\left(\max_{\pi\in\hat{\Pi}}|U(\pi,D)-U(\pi,P)|\geq\sqrt{\frac{\ln(2|\hat{\Pi}|/\delta)}{2T}}\right)\leq\delta.$$

Example: ϵ -net.

In a n dimensional space we require $|\hat{\Pi}| = O(\epsilon^{-n})$. This means that our statistical error is $O(\sqrt{n \ln(1/\epsilon\delta)/T})$.

 $^{^{1}}$ See Hoeffding's inequality in the confidence intervals presentation $\stackrel{>}{=}$ $\stackrel{>}{=}$

The finite hypothesis algorithm

- ▶ Input: a finite set of rules $\hat{\Pi}$, data D, utility U
- ▶ Return $\hat{\pi} \in \arg\max_{\pi \in \hat{\Pi}} U(\pi, D)$.

Regret of the finite hypothesis algorithm.

With probability $1-\delta$

$$U(\hat{\pi}, P) \ge U(\hat{\pi}^*, P) - \sqrt{2 \ln(2|\hat{\Pi}|/\delta)/T}$$
 (1)

$$U(\pi^*, P) - U(\hat{\pi}, P) \le \Delta + \sqrt{2\ln(2|\hat{H}|/\delta)/T}$$
 (2)

Examples

- ▶ ML estimation: $U(\theta, D) = P_{\theta}(D)$ is the data likelihood.
- ightharpoonup Accuracy, etc: $U(\pi, D)$.

VC Dimension

Here we consider sets Π of deterministic rules $\pi: X \to \{0,1\}$.

Shattering

If a $S \subset X$ can with |S| = m, can be assigned any labelling y_1, \ldots, y_m by a $\pi \in \Pi$, then we say Π shatters S.

The VC dimension

This is the largest-size set S that Π can shatter.

Example: Perceptrons on \mathbb{R}^2

This class has VC dimension 3 on the plane.

Binary classification

Learning algorithm λ

- ▶ Takes data $D = \{(x_t, y_t)\}$ as input
- ▶ Generates deterministic decision rules $\pi: X \to \{0, 1\}$,

The loss of a rule π .

- Assume an existing concept class $\pi^* \in \Pi$
- ▶ Distribution $x_t \sim P$ is i.i.d. and $x_1, \ldots, x_T \sim P^T$.
- ► The loss under distribution *P* is

$$L(\pi) = P(\{x : \pi(x) \neq \pi^*(x)\})$$

Realisable PAC learner

▶ $\lambda : (X \times Y)^* \to \Pi$ is (ϵ, δ) -PAC, if for any P and $\epsilon, \delta > 0$, and any concept $\pi^* \in \Pi$, there is T such that

$$P^{T}(\{D: L[\lambda(D)] > \epsilon\}) < \delta, \qquad D = (\{x_{t}, \pi^{*}(x_{t})\}), x_{t} \sim P.$$