Machine learning Part 2

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Recap of part 1

Supervised learning

- Regression: least-squares linear regression
- Classification: logistic regression

Regularization

• ℓ_2 a.k.a. ridge regularization

Model evaluation and selection

- Out-of-sample generalization; independent test set
- · Performance metrics:
 - regression: mean squared error
 - · classification: accuracy, ROC curve
- Cross-validation

Don't remember? watch Part 1 again!

Notation & vocabulary Supervised learning framework

$$Y = f(X) + E$$

• $Y \in \mathbb{R}$: output (a.k.a. target, dependent variable) to predict • $X \in \mathbb{R}^p$: features (a.k.a. inputs, regressors, descriptors,

- independent variables)
- $E \in \mathbb{R}$: unmodelled noise
- f: the function we try to approximate

Example: linear regression

$$Y = \beta_0 + \langle X, \beta \rangle + E$$

$$\frac{p}{\sqrt{y}}$$

$$= \beta_0 + \langle X, \beta \rangle + E$$

$$= \beta_0 + \sum_{j=1}^p X_j \beta_j + E$$

(2)

(1)

"learning" = estimating $\beta_0 \in \mathbb{R}$ and $\beta \in \mathbb{R}^p$

Dimensionality reduction

Until now



Add a step in the pipeline: simplifying the inputs



Dimensionality reduction

Problems when the number of features p becomes large

- Bigger errors on test data (larger variance of predictions)
- · Numerical stability issues
- · Computational cost and memory usage

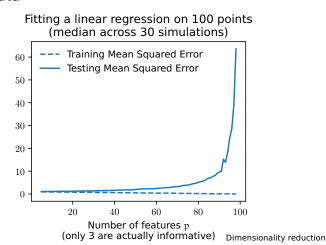
Toy example: linear regression with simulated data

- Generate $X \in \mathbb{R}^{n \times 3}$, $\beta \in \mathbb{R}^3$, and $y = X \beta \in \mathbb{R}^n$
- Append columns containing random noise to X
- Now $X \in \mathbb{R}^{n \times p}$, with $p \geqslant 3$, but only the first 3 columns are linked with y
- Split into training and testing tests and evaluate a linear regression model: what happens when p becomes large?

See sklearn.datasets.make_regression for generating data

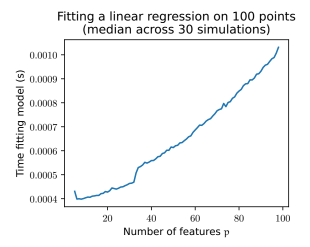
Model complexity: overfitting

- · Model complexity increases with dimension.
- Example: a linear model in dimension $\mathfrak p$ can fit exactly (0 training error) any set of $\mathfrak p+1$ points.
- Risk of overfitting: fitting exactly training data but failing on test data



Cost of fitting many parameters

- Many algorithms require polynomial time in p
- Implementations often make copies of the design matrix (e.g. for centering & rescaling)



Univariate feature selection

- a.k.a. feature screening, filtering . . .
- Check features (columns of X) one by one for association with the output y
- Keep only a fixed number or percentage of the features

Simple (linear) association criteria

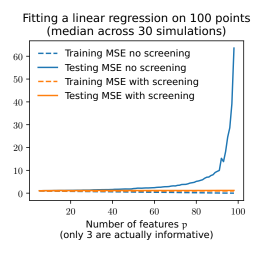
- for regression: correlation
- for classification: ANalysis Of VAriance

Read more in the scikit-learn user guide

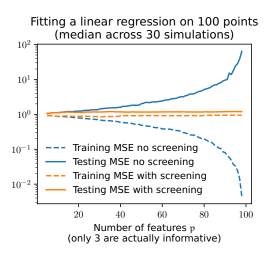
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https://scikit-learn.org/stable/modules/feature_selection.html#feature-selection
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Univariate feature selection

Keeping only the 10 best features (most correlated with y)

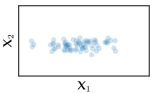


Same plot in log scale

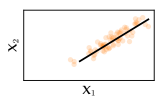


Linear decomposition methods

Maybe OK to drop X_2 :



Data low-dimensional but no feature can be dropped:

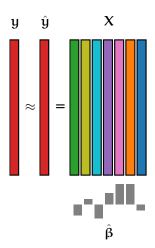


Find a better referential in which to represent the data

Linear regression: projection on the column

space of X

$$\hat{y} = X \hat{\beta}$$
 (4)

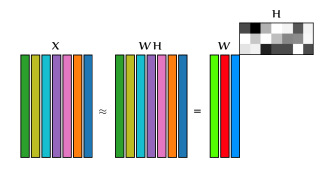


- Too many features: high variance & unstable solution
- Feature selection: drop some columns of X
- Other ways to build a family of k vectors on which to regress y?

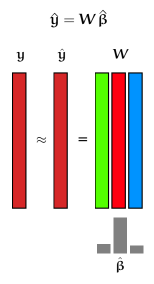
Linear decomposition: low-rank approximation of \mathbf{X}

Minimize

$$\|X - WH\|_{\mathsf{F}}^2 = \sum_{i,j} (X_{i,j} - (WH)_{i,j})^2$$
 (5)



Linear regression after dimensionality reduction



(6)

Prediction for a new data point $x \in \mathbb{R}^p$

- Find the combination of rows of H that is closest to x: regress x on H^T
- Multiply by $\hat{\beta}$

$$x \in \mathbb{R}^p o \mathsf{projection} o w \in \mathbb{R}^k o \langle \cdot \,, \, \hat{eta}
angle o \hat{\mathfrak{y}} \in \mathbb{R}$$
 (7)

Principal Component Analysis

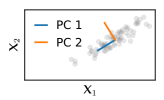
 Matrix factorization based on Singular Value Decomposition:

$$X = \mathbf{U} \, \mathbf{\Sigma} \, \mathbf{V}^{\mathsf{T}} \tag{8}$$

with $X \in \mathbb{R}^{n \times p}$, $U \in \mathbb{R}^{n \times p}$, $\Sigma \in \mathbb{R}^{p \times p}$, $V \in \mathbb{R}^{p \times p}$

- $\Sigma \succeq 0$ diagonal with decreasing values $\Sigma_{j,j}$ along the diagonal
- $\mathbf{U}^\mathsf{T} \mathbf{U} = \mathbf{I}_\mathsf{p}$
- $\mathbf{V}^{\mathsf{T}}\mathbf{V} = \mathbf{I}_{\mathsf{p}}$

Truncating the SVD to keep only the first ${\tt k}$ components gives the best rank- ${\tt k}$ approximation of X



Other decomposition methods

Many other methods use the same objective (sum of squared reconstruction errors), but add penalties or constraints on the factors

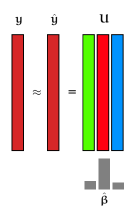
- Dictionary Learning
- · Non-negative Matrix Factorization
- · K-means clustering
- ..

What about u?

- PCA is an example of unsupervised learning: it does not use y
- Some other methods take it into account: e.g. Partial Least Squares

Ridge regression and PCA

- Both ridge regression and PC regression compute the coordinates of y in the basis given by the SVD of X
- ridge shrinks the coefficients of Singular Vector j by a factor $\sigma_i^2/(\sigma_i^2+\lambda)$
- PC regression sets the coefficient to 0 for all but the k largest σ_{j}



Setting hyperparameters

How can we choose:

- Number of features or PCA components k?
- The ridge hyperparameter λ?

Try a few and pick the best one...
But measure its performance on separate data!

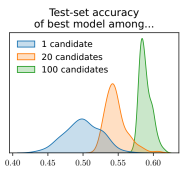
Need for fresh test data

When you hear "best", "maximum", "select", ... think "bias"

- I have 4 dice and want to find one that rolls high numbers
- I roll them all once and select the die that gives the highest number
- The selected die rolled a 5. Is 5 a good estimate of that die's average result? What if I had 1,000 dice?
- I need to roll it again to get an unbiased estimate

Accuracy of the best model

- Several models are trained, then evaluated on a separate test set
- All models give random answers expected accuracy is .5
- If I select the best one, its measured accuracy is biased . . .



Nested cross-validation

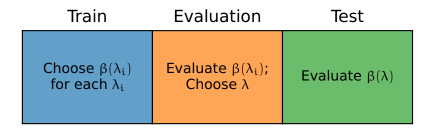
When you hear "best", "maximum", "select", ... think "bias" Setting the parameters

- Select β that gives the **best** prediction on training data
- The prediction score for $\hat{\beta}$ is biased: compute a new score on unseen test data.

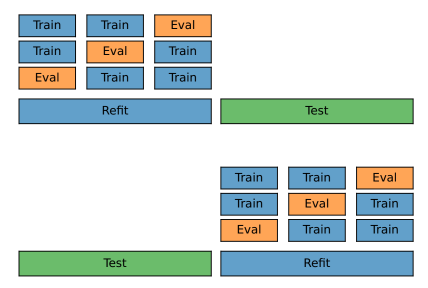
Setting the hyperparameters

- Repeat step 1 for a few values of λ, k, etc.., fitting and testing several models
- Select the hyperparameter that obtains the best prediction on test data
- The prediction score of that model on *test* data is biased: evaluate it again on unseen data

One split



Nested cross-validation



See sklearn.model_selection.GridSearchCV

Some common pitfalls with cross-validation

- Fitting part of the pipeline on the whole dataset
 - e.g. fit PCA on all data, then do cross-validation on dim-reduced dataset
 - USE sklearn.pipeline.Pipeline
- Ignoring dependencies between samples
 - · Multiple datapoints per participant
 - Time series
 - ...
- Ignoring dependencies between CV scores
 - Training sets overlap: cross-validation scores of different splits are not independent
- Over-interpreting good CV scores
 - Good CV scores on one dataset do not mean the model will always perform well on a new dataset

Supervised learning with fMRI

 Predict in which site / with which scanner a resting-state fMRI sequence was acquired

The decoding pipeline

- Masking: extracting voxels that are inside the brain
- Connectivity: measuring correlations between brain regions to build a feature vector for each participant
- Univariate feature selection with ANalysis Of VAriance
- Classifier: logistic regression

Implementation: in class